

Appendix E

Data Validation Reports



**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Boeing Realty Corp. C-6, EM2727
Collection Date: November 21, 2006
LDC Report Date: April 4, 2007
Matrix: Water
Parameters: Dissolved Gases
Validation Level: Tier 2 & 3
Laboratory: TestAmerica, Inc./Air Technology Laboratory, Inc.
Sample Delivery Group (SDG): IPK2470/A6112208-01/02

Sample Identification

IWC001_WG112106_0001**
MWC024_WG112106_0001

**Indicates sample underwent Tier 3 review

Introduction

This data review covers 2 water samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per Method RSK-175 for Dissolved Gases.

This review follows a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Organic Data Review (October 1999) as there are no current guidelines for the method stated above.

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

Blank results are summarized in Section III.

Field duplicates are summarized in Section IX.

Samples indicated by a double asterisk on the front cover underwent a Tier 3 review. Samples indicated by a single asterisk on the front cover underwent a Tier 2 review. Raw data were not evaluated for the samples reviewed by Tier 2 criteria since this review is based on QC data.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- N Presumptive evidence of presence of the constituent.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met with the following exceptions:

Sample	Compound	Total Days From Sample Collection Until Analysis	Required Holding Time (in Days) From Sample Collection Until Analysis	Flag	A or P
All samples in SDG IPK2470/A6112208-01/02	All TCL compounds	8	7	J (all detects) UJ (all non-detects)	P

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. Calibration

a. Initial Calibration

Initial calibration of compounds was performed as required by the method.

A curve fit, based on the initial calibration, was established for quantitation. The coefficient of determination (r^2) was greater than or equal to 0.990 .

b. Calibration Verification

Calibration verification was performed at required frequencies. The percent differences (%D) of amounts in continuing standard mixtures were within the 25.0% QC limits.

III. Blanks

Method blanks were reviewed for each matrix as applicable. No dissolved gas contaminants were found in the method blanks.

IV. Accuracy and Precision Data

a. Surrogate Recovery

Not required by the method.

b. Matrix Spike/Matrix Spike Duplicates

The laboratory has indicated that there were no matrix spike (MS) and matrix spike duplicate (MSD) analyses specified for the samples in this SDG, and therefore matrix spike and matrix spike duplicate analyses were not performed for this SDG.

c. Laboratory Control Samples

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

V. Target Compound Identification

All target compound identifications were within validation criteria on which a Tier 3 review was performed. Raw data were not evaluated for the samples reviewed by Tier 2 criteria.

VI. Compound Quantitation and CRQLs

All compound quantitation and CRQLs were within validation criteria on which a Tier 3 review was performed. Raw data were not evaluated for the samples reviewed by Tier 2 criteria.

VII. System Performance

The system performance was acceptable for samples on which a Tier 3 review was performed. Raw data were not evaluated for the samples reviewed by Tier 2 criteria.

VIII. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

IX. Field Duplicates

No field duplicates were identified in this SDG.

X. Field Blanks

No field blanks were identified in this SDG.

Boeing Realty Corp. C-6, EM2727

Dissolved Gases - Data Qualification Summary - SDG IPK2470/A6112208-01/02

SDG	Sample	Compound	Flag	A or P	Reason
IPK2470/ A6112208-01/02	IWC001_WG112106_0001** MWC024_WG112106_0001	All TCL compounds	J (all detects) UJ (all non-detects)	P	Technical holding times

Boeing Realty Corp. C-6, EM2727

Dissolved Gases - Laboratory Blank Data Qualification Summary - SDG IPK2470/A6112208-01/02

No Sample Data Qualified in this SDG

Client: TestAmerica
 Attn: Michele Chamberlin

Page 2 of 3
 A6112208

Client's Project: IPK2470
 Date Received: 11/22/2006
 Matrix: Water
 Units: ug/L

Dissolved Gases by EPA Procedure RSKSOP-175					
Lab No.:		A6112208-01	A6112208-02		
Client Sample I.D.:		IPK2470-07	IPK2470-09		
Date Sampled:		11/21/2006	11/21/2006		
Date Analyzed:		11/29/2006	11/29/2006		
Analyst Initials:		DT	DT		
Data File:		29nov013	29nov014		
QC Batch:		061129GCRA1	061129GCRA1		
Dilution Factor:		1.0	1.0		
ANALYTE	PQL	RL	Results	RL	Results
Methane	1.0	1.0	9.5 J	1.0	1,700 J
Ethane	2.0	2.0	ND UJ	2.0	ND UJ
Ethylene	3.0	3.0	ND J	3.0	ND J
Carbon Dioxide	200	200	16,000 J	200	17,000 J
Nitrogen	1,500	1,500	31,000 J	1,500	26,000 J

PQL = Practical Quantitation Limit
 ND = Not Detected (Below RL)
 RL = PQL X Dilution Factor

Reviewed/Approved By: Mark J. Johnson
 Mark J. Johnson
 Operations Manager

Date: 12-4-06

The cover letter is an integral part of this analytical report.

Handwritten signature/initials



AirTECHNOLOGY Laboratories, Inc.

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3/109

METHOD: GC Dissolved Gases (Method RSK-175)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	SW	Sampling dates: <u>11/29/06</u>
IIa.	Initial calibration	A	<u>1² 20-990</u>
IIb.	Calibration verification	A	<u>CCV ≤ 25</u>
III.	Blanks	A	
IVa.	Surrogate recovery	N	<u>not required</u>
IVb.	Matrix spike/Matrix spike duplicates	N	<u>client specified</u>
IVc.	Laboratory control samples	A	<u>LCSD</u>
V.	Target compound identification	A	Not reviewed for Tier II validation.
VI.	Compound Quantitation and CRQLs	A	Not reviewed for Tier II validation.
VII.	System Performance	A	Not reviewed for Tier II validation.
VIII.	Overall assessment of data	A	
IX.	Field duplicates	N	
X.	Field blanks	N	

Note: A = Acceptable ND = No compounds detected D = Duplicate
 N = Not provided/applicable R = Rinsate TB = Trip blank
 SW = See worksheet FB = Field blank EB = Equipment blank

Validated Samples: ** Indicates sample underwent Tier III validation

Water

1 [†]	IWC001_WG112106_0001**	11	<u>MS - 11/29/06</u>	21		31	
2 [†]	MWC024_WG112106_0001	12		22		32	
3		13		23		33	
4		14		24		34	
5		15		25		35	
6		16		26		36	
7		17		27		37	
8		18		28		38	
9		19		29		39	
10		20		30		40	

Notes: _____

LDC #: 16470AS)
 SDG #: see cover

VALIDATION FINDINGS CHECKLIST

Page: 1 of 2
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

Method: GC HPLC

Validation Area	Yes	No	NA	Findings/Comments
I. Technical holding times				
All technical holding times were met.		✓		
Cooler temperature criteria was met.	✓			
II. Initial calibration				
Did the laboratory perform a 5 point calibration prior to sample analysis?	✓			
Was a linear fit used for evaluation? If yes, were all percent relative standard deviations (%RSD) < 20%?		✓		
Was a curve fit used for evaluation? If Yes, what was the acceptance criteria used?	✓			
Did the initial calibration meet the curve fit acceptance criteria?	✓			
Were the RT windows properly established?	✓			
IV. Continuing calibration				
What type of continuing calibration calculation was performed? ___%D or %R	✓			
Was a continuing calibration analyzed daily?	✓			
Were all percent differences (%D) < 15%.0 or percent recoveries 85-115%?	✓			
Were all the retention times within the acceptance windows?	✓			
V. Blanks				
Was a method blank associated with every sample in this SDG?	✓			
Was a method blank analyzed for each matrix and concentration?	✓			
Was there contamination in the method blanks? If yes, please see the Blanks validation completeness worksheet.			✓	
VI. Surrogate spikes				
Were all surrogate %R within the QC limits?			✓	
If the percent recovery (%R) of one or more surrogates was outside QC limits, was a reanalysis performed to confirm %R?			✓	
If any %R was less than 10 percent, was a reanalysis performed to confirm %R?			✓	
VII. Matrix spike/Matrix spike duplicates				
Were a matrix spike (MS) and matrix spike duplicate (MSD) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD. Soil / Water.			✓	
Was a MS/MSD analyzed every 20 samples of each matrix?			✓	
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the QC limits?			✓	
VIII. Laboratory control samples				
Was an LCS analyzed for this SDG?	✓			
Was an LCS analyzed per extraction batch?	✓			

DC #: 16470AS1
 JG #: see cover

VALIDATION FINDINGS CHECKLIST

Page: 2 of 2
 Reviewer: FA
 2nd Reviewer: RA

Validation Area	Yes	No	NA	Findings/Comments
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the QC limits? <i>XII. Regional Quality Assurance and Quality Control</i>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were performance evaluation (PE) samples performed? <i>XIII. Laboratory Validation</i>	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were the performance evaluation (PE) samples within the acceptance limits? <i>XIII. Laboratory Validation</i>	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were the retention times of reported detects within the RT windows? <i>XIII. Laboratory Validation</i>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were compound quantitation and CRQLs adjusted to reflect all sample dilutions and dry weight factors applicable to level IV validation? <i>XIII. Laboratory Validation</i>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
System performance was found to be acceptable. <i>XIII. Laboratory Validation</i>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Overall assessment of data was found to be acceptable. <i>XIII. Laboratory Validation</i>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were field duplicate pairs identified in this SDG? <i>XIV. Field Duplicates</i>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
Were target compounds detected in the field duplicates? <i>XIV. Field Duplicates</i>	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were field blanks identified in this SDG? <i>XV. Field Blanks</i>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
Were target compounds detected in the field blanks? <i>XV. Field Blanks</i>	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	

LDC# 16470As1
 SDG# pu ever

VALIDATION FINDINGS WORKSHEET
 Initial Calibration Calculation Verification

Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: A

METHOD: RSK-175

Calibration Date	Column/ Detector	Compound	Standard	X	Y
05/23/06	FID (middle)	methane	Point 1	10	9309
			Point 2	100	89744
			Point 3	1000	863867
			Point 4	5000	4401745
			Point 5	10000	9597354
			Point 6		
			Point 7		
			Point 8		

Regression Output:	Recalculated Result	Result Reported by the Laboratory
Constant	0	0
Std Err of Y Est	181942.83552443456	
R Squared	0.99804	0.998417
No. of Observations	5	
Degrees of Freedom	4	
X Coefficient(s)	943.21962676	9.4322E+02
Std Err of Coef.	16.2081064622	

LDC #: 16470AS1
 SDG #: per cover

VALIDATION FINDINGS WORKSHEET
 Laboratory Control Sample/Laboratory Control Sample Duplicate Results Verification

Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: GC HPLC

The percent recoveries (%R) and relative percent differences (RPD) of the laboratory control sample and laboratory control sample duplicate were recalculated for the compounds identified below using the following calculation:

%Recovery = $100 * ((SSC - SC) / SA)$ Where SSC = Spiked concentration
 SA = Spike added SC = Sample concentration

RPD = $((|SSCLCS - SSCLCSD| * 2) / (SSCLCS + SSCLCSD)) * 100$ LCS = Laboratory Control Sample percent recovery
 LCSD = Laboratory Control Sample duplicate percent recovery

LCS/LCSD samples: LCS 1/D

Compound	Spike Added (ug/L)		Sample Conc. (ug/L)		Spike Sample Concentration (ug/L)		LCS		LCSD		LCS		LCSD		LCS/LCSD		
	LCS	LCSD	LCS	LCSD	LCS	LCSD	Reported	Recalc.	Reported	Recalc.	Reported	Recalc.	Reported	Recalc.	Reported	Recalc.	
							Percent Recovery		Percent Recovery		Percent Recovery		Percent Recovery		RPD		
Gasoline (8015)																	
Diesel (8015)																	
Benzene (8021B)																	
Methane (F8K-175)	6900	6900	0	0	7694.7	7376.5	111	111.5	106	106.9	4.2	4.2					
2,4-D (8151)																	
Dinoseb (8151)																	
Naphthalene (8310)																	
Anthracene (8310)																	
HMX (8330)																	
2,4,6-Trinitrotoluene (8330)																	

Comments: Refer to Laboratory Control Sample/Laboratory Control Sample Duplicate findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 16470A57
 SDG #: all covered

VALIDATION FINDINGS WORKSHEET
Sample Calculation Verification

Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: GC HPLC

Y N N/A
 Y N N/A

Were all reported results recalculated and verified for all level IV samples?
 Were all recalculated results for detected target compounds within 10% of the reported results?

Concentration = $\frac{(A)(Fv)(Df)}{(RF)(Vs \text{ or } Ws)(\%S/100)}$

- A= Area or height of the compound to be measured
- Fv= Final Volume of extract
- Df= Dilution Factor
- RF= Average response factor of the compound in the initial calibration
- Vs= Initial volume of the sample
- Ws= Initial weight of the sample
- %S= Percent Solid

Example: Sample ID: #1 Compound Name: Methane

Concentration = _____

$$y = 9.4322 \times 10^2 (x)$$

$$94667 = 9.4322 \times 10^2 (x)$$

$$x = 100.4$$

#	Sample ID	Compound	Reported Concentrations	Recalculated Results Concentrations	Qualifications
		gas in HS	$(100.4)(55.51)$ $(1000)(41300)$	$(16.04)(1000)$ $(1000)(41300)$	2.1645
		gas in liquid	$(100.4)(16.04)$ $(1000)(22.4)$	$(1000)(4)$ $(36)(278)$ (273)	7.318
				Total	9.48 ug/L

Comments: _____

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Boeing Realty Corp. C-6, EM2727
Collection Date: November 20, 2006
LDC Report Date: April 4, 2007
Matrix: Water
Parameters: Dissolved Gases
Validation Level: Tier 1
Laboratory: TestAmerica, Inc.

Sample Delivery Group (SDG): IPK2310

Sample Identification

MWC015_WG112006_0001

Introduction

This data review covers one water sample listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per Method RSK-175 for Dissolved Gases.

This review follows a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Organic Data Review (October 1999) as there are no current guidelines for the method stated above.

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified a P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

Blank results are summarized in Section III.

Field duplicates are summarized in Section IX.

Raw data were not reviewed for this SDG. The review was based on QC data.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- N Presumptive evidence of presence of the constituent.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. Calibration

a. Initial Calibration

Initial calibration data were not reviewed for Tier 1.

b. Calibration Verification

Calibration verification data were not reviewed for Tier 1.

III. Blanks

Method blanks were performed at the required frequency. No dissolved gas contaminants were found in the method blanks.

IV. Accuracy and Precision Data

a. Surrogate Recovery

Not required by the method.

b. Matrix Spike/Matrix Spike Duplicates

The laboratory has indicated that there were no matrix spike (MS) and matrix spike duplicate (MSD) analyses specified for the samples in this SDG, and therefore matrix spike and matrix spike duplicate analyses were not performed for this SDG.

c. Laboratory Control Samples

Laboratory control samples were analyzed at the required frequency. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

V. Target Compound Identification

Raw data were not reviewed for this SDG.

VI. Compound Quantitation and CRQLs

Raw data were not reviewed for this SDG.

VII. System Performance

Raw data were not reviewed for this SDG.

VIII. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

IX. Field Duplicates

No field duplicates were identified in this SDG.

X. Field Blanks

No field blanks were identified in this SDG.

**Boeing Realty Corp. C-6, EM2727
Dissolved Gases - Data Qualification Summary - SDG IPK2310**

No Sample Data Qualified in this SDG

**Boeing Realty Corp. C-6, EM2727
Dissolved Gases - Laboratory Blank Data Qualification Summary - SDG IPK2310**

No Sample Data Qualified in this SDG

TestAmerica

ANALYTICAL TESTING CORPORATION

TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IPK2310

Sampled: 11/20/06
Received: 11/20/06

RSK175 Dissolved Gases in Water

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IPK2310-05 (MWC015_WG112006_0001 - Water) - cont.									
Reporting Units: ug/L									
Methane	RSK-175	361129GC8A	0.39	1.0	10	1	11/29/06	11/29/06	
Ethane	RSK-175	361129GC8A	0.50	2.0	ND	1	11/29/06	11/29/06	
Ethylene	RSK-175	361129GC8A	0.33	3.0	ND	1	11/29/06	11/29/06	
Nitrogen	RSK-175	361129GC8A	222	1500	25000	1	11/29/06	11/29/06	

TestAmerica - Irvine, CA
Nicholas Marz For Michele Chamberlin
Project Manager

11/20/06

The results pertain only to the samples tested in the laboratory. This report shall not be reproduced, except in full, without written permission from TestAmerica.

IPK2310 <Page 20 of 44>

METHOD: GC Dissolved Gases (Method RSK-175)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: 11/20/06
IIa.	Initial calibration	N	
IIb.	Calibration verification	N	
III.	Blanks	A	
IVa.	Surrogate recovery	N	not required
IVb.	Matrix spike/Matrix spike duplicates	N	client specified
IVc.	Laboratory control samples	A	LCBID
V.	Target compound identification	N	
VI.	Compound Quantitation and CRQLs	N	
VII.	System Performance	N	
VIII.	Overall assessment of data	A	
IX.	Field duplicates	N	
X.	Field blanks	N	

Note: A = Acceptable ND = No compounds detected D = Duplicate
 N = Not provided/applicable R = Rinsate TB = Trip blank
 SW = See worksheet FB = Field blank EB = Equipment blank

Validated Samples: water

1	MWC015_WG112006_0001	11	MB - 11/29/06	21		31	
2		12		22		32	
3		13		23		33	
4		14		24		34	
5		15		25		35	
6		16		26		36	
7		17		27		37	
8		18		28		38	
9		19		29		39	
10		20		30		40	

Notes: _____

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Boeing Realty Corp. C-6, EM2727
Collection Date: November 20, 2006
LDC Report Date: March 28, 2007
Matrix: Water
Parameters: Dissolved Manganese
Validation Level: Tier 1
Laboratory: TestAmerica

Sample Delivery Group (SDG): IPK2310

Sample Identification

MWC015_WG112006_0001
MWC015_WG112006_0001MS
MWC015_WG112006_0001MSD

Introduction

This data review covers 3 water samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 6010B for Dissolved Manganese.

This review follows a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Inorganic Data Review (October 2004) as there are no current guidelines for the methods stated above.

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

Blanks are summarized in Section III.

Field duplicates are summarized in Section XIII.

Samples indicated by a double asterisk on the front cover underwent a Tier 2 review. A Tier 1 review was performed on all of the other samples. Raw data were not evaluated for the samples reviewed by Tier 1 and 2 criteria since this review is based on QC data.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- N Presumptive evidence of presence of the constituent.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. Calibration

Calibration data were not reviewed for Tier 1.

III. Blanks

Method blanks were reviewed for each matrix as applicable. No contaminant concentrations were found in the preparation blanks.

IV. ICP Interference Check Sample (ICS) Analysis

ICP interference check sample analysis data were not reviewed for Tier 1.

V. Matrix Spike Analysis

Matrix spike (MS) and matrix spike duplicate (MSD) samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

VI. Duplicate Sample Analysis

Duplicate (DUP) sample analyses were reviewed for each matrix as applicable.

VII. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) were within QC limits.

VIII. Internal Standards

ICP-MS was not utilized in this SDG.

IX. Furnace Atomic Absorption QC

Graphite furnace atomic absorption was not utilized in this SDG.

X. ICP Serial Dilution

ICP serial dilution was not performed for this SDG.

XI. Sample Result Verification

Raw data were not reviewed for this SDG.

XII. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

XIII. Field Duplicates

No field duplicates were identified in this SDG.

XIV. Field Blanks

No field blanks were identified in this SDG.

**Boeing Realty Corp. C-6, EM2727
Dissolved Manganese - Data Qualification Summary - SDG IPK2310**

No Sample Data Qualified in this SDG

**Boeing Realty Corp. C-6, EM2727
Dissolved Manganese - Laboratory Blank Data Qualification Summary - SDG
IPK2310**

No Sample Data Qualified in this SDG

TestAmerica

ANALYTICAL TESTING CORPORATION

TAT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IPK2310

Sampled: 11/20/06
Received: 11/20/06

DISSOLVED METALS

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IPK2310-05 (MWC015_WG112006_0001 - Water)									
Reporting Units: mg/l									
Manganese	EPA 6010B-Diss	6K20133	N/A	0.020	0.022	1	11/20/06	11/20/06	

TestAmerica - Irvine, CA
Nicholas Marz For Michele Chamberlin
Project Manager

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LDC #: 16470B4
 SDG #: IPK2310
 Laboratory: Test America

VALIDATION COMPLETENESS WORKSHEET
 EPA Region 1 - Tier 1

Date: 3/27/07
 Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: Dissolved Manganese (EPA SW 846 Method 6010B)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: <u>11/20/06</u>
II.	Calibration	N	
III.	Blanks	A	
IV.	ICP Interference Check Sample (ICS) Analysis	N	
V.	Matrix Spike Analysis	A	<u>3 runs MSD</u>
VI.	Duplicate Sample Analysis	N	
VII.	Laboratory Control Samples (LCS)	A	<u>LCS</u>
VIII.	Internal Standard (ICP-MS)	N	<u>} not utilized</u>
IX.	Furnace Atomic Absorption QC	N	
X.	ICP Serial Dilution	N	<u>not performed.</u>
XI.	Sample Result Verification	N	
XII.	Overall Assessment of Data	A	
XIII.	Field Duplicates	N	
XIV.	Field Blanks	N	

Note: A = Acceptable
 N = Not provided/applicable
 SW = See worksheet
 ND = No compounds detected
 R = Rinsate
 FB = Field blank
 D = Duplicate
 TB = Trip blank
 EB = Equipment blank

Validated Samples: A2

1	MWC015_WG112006_0001	11		21		31	
2	MWC015_WG112006_0001MS	12		22		32	
3	MWC015_WG112006_0001MSD	13		23		33	
4	<u>PB</u>	14		24		34	
5		15		25		35	
6		16		26		36	
7		17		27		37	
8		18		28		38	
9		19		29		39	
10		20		30		40	

Notes: _____

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Boeing Realty Corp. C-6, EM2727
Collection Date: November 20, 2006
LDC Report Date: April 5, 2007
Matrix: Water
Parameters: Volatiles
Validation Level: Tier 1
Laboratory: TestAmerica

Sample Delivery Group (SDG): IPK2310

Sample Identification

MWC015_WG112006_0001
MWC015_WG112006_0001RE1
MWC016_WG112006_0001
MWC016_WG112006_0001RE1
MWC015_WG112006_0001MS
MWC015_WG112006_0001MSD

2

Introduction

This data review covers 6 water samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 8260B for Volatiles.

This review follows a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Organic Data Review (October 1999) as there are no current guidelines for the method stated above.

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

Blank results are summarized in Section V.

Field duplicates are summarized in Section XVI.

Raw data were not reviewed for this SDG. The review was based on QC data.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- N Presumptive evidence of presence of the constituent.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. GC/MS Instrument Performance Check

Instrument performance data were not reviewed for Tier 1.

III. Initial Calibration

Initial calibration data were not reviewed for Tier 1.

IV. Continuing Calibration

Continuing calibration data were not reviewed for Tier 1.

V. Blanks

Method blanks were reviewed for each matrix as applicable. No volatile contaminants were found in the method blanks with the following exceptions:

Method Blank ID	Analysis Date	Compound TIC (RT in minutes)	Concentration	Associated Samples
6K22012-BLK	11/22/06	Tetrahydrofuran	4.41 ug/L	All samples in SDG IPK2310

Sample concentrations were compared to concentrations detected in the method blanks. The sample concentrations were either not detected or were significantly greater (>10X for common contaminants, >5X for other contaminants) than the concentrations found in the associated method blanks.

VI. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VII. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the following exceptions:

Spike ID (Associated Samples)	Compound	MS (%R) (Limits)	MSD (%R) (Limits)	RPD (Limits)	Flag	A or P
MWC015_WG112006_0001MS/MSD (MWC015_WG112006_0001)	Bromodichloromethane	-	-	38 (≤ 20)	J (all detects) UJ (all non-detects)	A

VIII. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) were within QC limits.

IX. Regional Quality Assurance and Quality Control

Not applicable.

X. Internal Standards

Internal standards data were not reviewed for Tier 1.

XI. Target Compound Identifications

Raw data were not reviewed for this SDG.

XII. Compound Quantitation and CRQLs

Raw data were not reviewed for this SDG.

XIII. Tentatively Identified Compounds (TICs)

Raw data were not reviewed for this SDG.

XIV. System Performance

Raw data were not reviewed for this SDG.

XV. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

XVI. Field Duplicates

No field duplicates were identified in this SDG.

XVII. Field Blanks

No field blanks were identified in this SDG.

Boeing Realty Corp. C-6, EM2727
Volatiles - Data Qualification Summary - SDG IPK2310

SDG	Sample	Compound	Flag	A or P	Reason
IPK2310	MWC015_WG112006_0001	Bromodichloromethane	J (all detects) UJ (all non-detects)	A	Matrix spike/Matrix spike duplicates (RPD)

Boeing Realty Corp. C-6, EM2727
Volatiles - Laboratory Blank Data Qualification Summary - SDG IPK2310

No Sample Data Qualified in this SDG

TestAmerica

ANALYTICAL TESTING CORPORATION

TAIT Environmental/Boeing 701 N. Parkcenter Drive Santa Ana, CA 92705 Attention: Mehmet Pehlivan	Project ID: Boeing C-6 Torrance EM2727 Report Number: IPK2310	Sampled: 11/20/06 Received: 11/20/06
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VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IPK2310-05 (MWC015_WG112006_0001 - Water)									
Reporting Units: ug/l									
Acetone	EPA 8260B	6K22012	4.5	10	ND	1	11/22/06	11/22/06	
Benzene	EPA 8260B	6K22012	0.28	1.0	ND	1	11/22/06	11/22/06	
Bromobenzene	EPA 8260B	6K22012	0.27	1.0	ND	1	11/22/06	11/22/06	
Bromochloromethane	EPA 8260B	6K22012	0.32	1.0	ND	1	11/22/06	11/22/06	
Bromodichloromethane	EPA 8260B	6K22012	0.30	1.0	ND	1	11/22/06	11/22/06	R
Bromoform	EPA 8260B	6K22012	0.40	1.0	ND	1	11/22/06	11/22/06	
Bromomethane	EPA 8260B	6K22012	0.42	1.0	ND	1	11/22/06	11/22/06	
2-Butanone (MEK)	EPA 8260B	6K22012	3.8	5.0	ND	1	11/22/06	11/22/06	
n-Butylbenzene	EPA 8260B	6K22012	0.37	1.0	ND	1	11/22/06	11/22/06	
sec-Butylbenzene	EPA 8260B	6K22012	0.25	1.0	ND	1	11/22/06	11/22/06	
tert-Butylbenzene	EPA 8260B	6K22012	0.22	1.0	ND	1	11/22/06	11/22/06	
Carbon Disulfide	EPA 8260B	6K22012	0.48	1.0	ND	1	11/22/06	11/22/06	
Carbon tetrachloride	EPA 8260B	6K22012	0.28	0.50	ND	1	11/22/06	11/22/06	
Chlorobenzene	EPA 8260B	6K22012	0.36	1.0	ND	1	11/22/06	11/22/06	
Chloroethane	EPA 8260B	6K22012	0.40	2.0	ND	1	11/22/06	11/22/06	
Chloroform	EPA 8260B	6K22012	0.33	1.0	1.8	1	11/22/06	11/22/06	
Chloromethane	EPA 8260B	6K22012	0.40	2.0	ND	1	11/22/06	11/22/06	
2-Chlorotoluene	EPA 8260B	6K22012	0.28	1.0	ND	1	11/22/06	11/22/06	
4-Chlorotoluene	EPA 8260B	6K22012	0.29	1.0	ND	1	11/22/06	11/22/06	
1,2-Dibromo-3-chloropropane	EPA 8260B	6K22012	0.97	2.0	ND	1	11/22/06	11/22/06	
Dibromochloromethane	EPA 8260B	6K22012	0.28	1.0	ND	1	11/22/06	11/22/06	
1,2-Dibromoethane (EDB)	EPA 8260B	6K22012	0.40	1.0	ND	1	11/22/06	11/22/06	
1,4-Dichlorobenzene	EPA 8260B	6K22012	0.37	1.0	ND	1	11/22/06	11/22/06	
1,2-Dichlorobenzene	EPA 8260B	6K22012	0.32	1.0	ND	1	11/22/06	11/22/06	
1,3-Dichlorobenzene	EPA 8260B	6K22012	0.35	1.0	ND	1	11/22/06	11/22/06	
Dichlorodifluoromethane	EPA 8260B	6K22012	0.79	1.0	ND	1	11/22/06	11/22/06	
1,2-Dichloroethane	EPA 8260B	6K22012	0.28	0.50	ND	1	11/22/06	11/22/06	
1,1-Dichloroethane	EPA 8260B	6K22012	0.27	1.0	ND	1	11/22/06	11/22/06	
1,1-Dichloroethene	EPA 8260B	6K22012	0.42	1.0	4.6	1	11/22/06	11/22/06	
cis-1,2-Dichloroethene	EPA 8260B	6K22012	0.32	1.0	4.6	1	11/22/06	11/22/06	
trans-1,2-Dichloroethene	EPA 8260B	6K22012	0.27	1.0	ND	1	11/22/06	11/22/06	
1,2-Dichloropropane	EPA 8260B	6K22012	0.35	1.0	ND	1	11/22/06	11/22/06	
2,2-Dichloropropane	EPA 8260B	6K22012	0.34	1.0	ND	1	11/22/06	11/22/06	
cis-1,3-Dichloropropene	EPA 8260B	6K22012	0.22	0.50	ND	1	11/22/06	11/22/06	
1,1-Dichloropropene	EPA 8260B	6K22012	0.28	1.0	ND	1	11/22/06	11/22/06	
trans-1,3-Dichloropropene	EPA 8260B	6K22012	0.32	0.50	ND	1	11/22/06	11/22/06	
Ethylbenzene	EPA 8260B	6K22012	0.25	1.0	ND	1	11/22/06	11/22/06	
Hexachlorobutadiene	EPA 8260B	6K22012	0.38	1.0	ND	1	11/22/06	11/22/06	
2-Hexanone	EPA 8260B	6K22012	2.6	6.0	ND	1	11/22/06	11/22/06	
Iodomethane	EPA 8260B	6K22012	1.0	2.0	ND	1	11/22/06	11/22/06	C
Isopropylbenzene	EPA 8260B	6K22012	0.25	1.0	ND	1	11/22/06	11/22/06	

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Nicholas Marz For Michele Chamberlin
Project Manager

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ANALYTICAL TESTING CORPORATION

TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IPK2310

Sampled: 11/20/06
Received: 11/20/06

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IPK2310-05 (MWC015_WG112006_0001 - Water) - cont.									
Reporting Units: ug/l									
p-Isopropyltoluene	EPA 8260B	6K22012	0.28	1.0	ND	1	11/22/06	11/22/06	
Methyl-tert-butyl Ether (MTBE)	EPA 8260B	6K22012	0.32	1.0	ND	1	11/22/06	11/22/06	
Methylene chloride	EPA 8260B	6K22012	0.95	1.0	ND	1	11/22/06	11/22/06	
4-Methyl-2-pentanone (MIBK)	EPA 8260B	6K22012	3.5	5.0	ND	1	11/22/06	11/22/06	
n-Propylbenzene	EPA 8260B	6K22012	0.27	1.0	ND	1	11/22/06	11/22/06	
Styrene	EPA 8260B	6K22012	0.16	1.0	ND	1	11/22/06	11/22/06	
1,1,1,2-Tetrachloroethane	EPA 8260B	6K22012	0.27	1.0	ND	1	11/22/06	11/22/06	
1,1,2,2-Tetrachloroethane	EPA 8260B	6K22012	0.24	1.0	ND	1	11/22/06	11/22/06	
Tetrachloroethene	EPA 8260B	6K22012	0.32	1.0	ND	1	11/22/06	11/22/06	
Tetrahydrofuran (THF)	EPA 8260B	6K22012	3.5	10	ND	1	11/22/06	11/22/06	
Toluene	EPA 8260B	6K22012	0.36	1.0	ND	1	11/22/06	11/22/06	
1,2,3-Trichlorobenzene	EPA 8260B	6K22012	0.30	1.0	ND	1	11/22/06	11/22/06	
1,2,4-Trichlorobenzene	EPA 8260B	6K22012	0.48	1.0	ND	1	11/22/06	11/22/06	
1,1,2-Trichloroethane	EPA 8260B	6K22012	0.30	1.0	ND	1	11/22/06	11/22/06	
1,1,1-Trichloroethane	EPA 8260B	6K22012	0.30	1.0	ND	1	11/22/06	11/22/06	
Trichlorofluoromethane	EPA 8260B	6K22012	0.34	2.0	ND	1	11/22/06	11/22/06	
1,2,3-Trichloropropane	EPA 8260B	6K22012	0.40	1.0	ND	1	11/22/06	11/22/06	
1,2,4-Trimethylbenzene	EPA 8260B	6K22012	0.23	1.0	ND	1	11/22/06	11/22/06	
1,3,5-Trimethylbenzene	EPA 8260B	6K22012	0.26	1.0	ND	1	11/22/06	11/22/06	
Vinyl acetate	EPA 8260B	6K22012	1.7	6.0	ND	1	11/22/06	11/22/06	
Vinyl chloride	EPA 8260B	6K22012	0.30	0.50	ND	1	11/22/06	11/22/06	
Xylenes, Total	EPA 8260B	6K22012	0.90	1.0	ND	1	11/22/06	11/22/06	
Surrogate: 4-Bromofluorobenzene (80-120%)					93 %				
Surrogate: Dibromofluoromethane (80-120%)					96 %				
Surrogate: Toluene-d8 (80-120%)					95 %				

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Project Manager

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ANALYTICAL TESTING CORPORATION

TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IPK2310

Sampled: 11/20/06
Received: 11/20/06

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IPK2310-05RE1 (MWC015_WG112006_0001 - Water) - cont.									
Reporting Units: ug/l									
Trichloroethene	EPA 8260B	6K22012	2.6	10	810	10	11/22/06	11/22/06	
Surrogate: 4-Bromofluorobenzene (80-120%)					94 %				
Surrogate: Dibromofluoromethane (80-120%)					100 %				
Surrogate: Toluene-d8 (80-120%)					98 %				

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Nicholas Marz For Michele Chamberlin
Project Manager

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TestAmerica

ANALYTICAL TESTING CORPORATION

TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IPK2310

Sampled: 11/20/06
Received: 11/20/06

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IPK2310-06 (MWC016_WG112006_0001 - Water)									
Reporting Units: ug/l									
Acetone	EPA 8260B	6K22012	4.5	10	ND	1	11/22/06	11/22/06	
Benzene	EPA 8260B	6K22012	0.28	1.0	ND	1	11/22/06	11/22/06	
Bromobenzene	EPA 8260B	6K22012	0.27	1.0	ND	1	11/22/06	11/22/06	
Bromochloromethane	EPA 8260B	6K22012	0.32	1.0	ND	1	11/22/06	11/22/06	
Bromodichloromethane	EPA 8260B	6K22012	0.30	1.0	ND	1	11/22/06	11/22/06	
Bromoform	EPA 8260B	6K22012	0.40	1.0	ND	1	11/22/06	11/22/06	
Bromomethane	EPA 8260B	6K22012	0.42	1.0	ND	1	11/22/06	11/22/06	
2-Butanone (MEK)	EPA 8260B	6K22012	3.8	5.0	ND	1	11/22/06	11/22/06	
n-Butylbenzene	EPA 8260B	6K22012	0.37	1.0	ND	1	11/22/06	11/22/06	
sec-Butylbenzene	EPA 8260B	6K22012	0.25	1.0	ND	1	11/22/06	11/22/06	
tert-Butylbenzene	EPA 8260B	6K22012	0.22	1.0	ND	1	11/22/06	11/22/06	
Carbon Disulfide	EPA 8260B	6K22012	0.48	1.0	ND	1	11/22/06	11/22/06	
Carbon tetrachloride	EPA 8260B	6K22012	0.28	0.50	1.0	1	11/22/06	11/22/06	
Chlorobenzene	EPA 8260B	6K22012	0.36	1.0	ND	1	11/22/06	11/22/06	
Chloroethane	EPA 8260B	6K22012	0.40	2.0	ND	1	11/22/06	11/22/06	
Chloroform	EPA 8260B	6K22012	0.33	1.0	8.8	1	11/22/06	11/22/06	
Chloromethane	EPA 8260B	6K22012	0.40	2.0	ND	1	11/22/06	11/22/06	
2-Chlorotoluene	EPA 8260B	6K22012	0.28	1.0	ND	1	11/22/06	11/22/06	
4-Chlorotoluene	EPA 8260B	6K22012	0.29	1.0	ND	1	11/22/06	11/22/06	
1,2-Dibromo-3-chloropropane	EPA 8260B	6K22012	0.97	2.0	ND	1	11/22/06	11/22/06	
Dibromochloromethane	EPA 8260B	6K22012	0.28	1.0	1.6	1	11/22/06	11/22/06	
1,2-Dibromoethane (EDB)	EPA 8260B	6K22012	0.40	1.0	ND	1	11/22/06	11/22/06	
1,4-Dichlorobenzene	EPA 8260B	6K22012	0.37	1.0	ND	1	11/22/06	11/22/06	
1,2-Dichlorobenzene	EPA 8260B	6K22012	0.32	1.0	ND	1	11/22/06	11/22/06	
1,3-Dichlorobenzene	EPA 8260B	6K22012	0.35	1.0	ND	1	11/22/06	11/22/06	
Dichlorodifluoromethane	EPA 8260B	6K22012	0.79	1.0	ND	1	11/22/06	11/22/06	
1,2-Dichloroethane	EPA 8260B	6K22012	0.28	0.50	ND	1	11/22/06	11/22/06	
1,1-Dichloroethane	EPA 8260B	6K22012	0.27	1.0	ND	1	11/22/06	11/22/06	
1,1-Dichloroethene	EPA 8260B	6K22012	0.42	1.0	7.8	1	11/22/06	11/22/06	
cis-1,2-Dichloroethene	EPA 8260B	6K22012	0.32	1.0	1.4	1	11/22/06	11/22/06	
trans-1,2-Dichloroethene	EPA 8260B	6K22012	0.27	1.0	ND	1	11/22/06	11/22/06	
1,2-Dichloropropane	EPA 8260B	6K22012	0.35	1.0	ND	1	11/22/06	11/22/06	
2,2-Dichloropropane	EPA 8260B	6K22012	0.34	1.0	ND	1	11/22/06	11/22/06	
cis-1,3-Dichloropropene	EPA 8260B	6K22012	0.22	0.50	ND	1	11/22/06	11/22/06	
1,1-Dichloropropene	EPA 8260B	6K22012	0.28	1.0	ND	1	11/22/06	11/22/06	
trans-1,3-Dichloropropene	EPA 8260B	6K22012	0.32	0.50	ND	1	11/22/06	11/22/06	
Ethylbenzene	EPA 8260B	6K22012	0.25	1.0	ND	1	11/22/06	11/22/06	
Hexachlorobutadiene	EPA 8260B	6K22012	0.38	1.0	ND	1	11/22/06	11/22/06	
2-Hexanone	EPA 8260B	6K22012	2.6	6.0	ND	1	11/22/06	11/22/06	
Iodomethane	EPA 8260B	6K22012	1.0	2.0	ND	1	11/22/06	11/22/06	C
Isopropylbenzene	EPA 8260B	6K22012	0.25	1.0	ND	1	11/22/06	11/22/06	

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Nicholas Marz For Michele Chamberlin
Project Manager

11/24/07

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ANALYTICAL TESTING CORPORATION

TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IPK2310

Sampled: 11/20/06
Received: 11/20/06

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IPK2310-06 (MWC016_WG112006_0001 - Water) - cont.									
Reporting Units: ug/l									
p-Isopropyltoluene	EPA 8260B	6K22012	0.28	1.0	ND	1	11/22/06	11/22/06	
Methyl-tert-butyl Ether (MTBE)	EPA 8260B	6K22012	0.32	1.0	ND	1	11/22/06	11/22/06	
Methylene chloride	EPA 8260B	6K22012	0.95	1.0	ND	1	11/22/06	11/22/06	
4-Methyl-2-pentanone (MIBK)	EPA 8260B	6K22012	3.5	5.0	ND	1	11/22/06	11/22/06	
n-Propylbenzene	EPA 8260B	6K22012	0.27	1.0	ND	1	11/22/06	11/22/06	
Styrene	EPA 8260B	6K22012	0.16	1.0	ND	1	11/22/06	11/22/06	
1,1,1,2-Tetrachloroethane	EPA 8260B	6K22012	0.27	1.0	ND	1	11/22/06	11/22/06	
1,1,2,2-Tetrachloroethane	EPA 8260B	6K22012	0.24	1.0	ND	1	11/22/06	11/22/06	
Tetrachloroethene	EPA 8260B	6K22012	0.32	1.0	ND	1	11/22/06	11/22/06	
Tetrahydrofuran (THF)	EPA 8260B	6K22012	3.5	10	ND	1	11/22/06	11/22/06	
Toluene	EPA 8260B	6K22012	0.36	1.0	ND	1	11/22/06	11/22/06	
1,2,3-Trichlorobenzene	EPA 8260B	6K22012	0.30	1.0	ND	1	11/22/06	11/22/06	
1,2,4-Trichlorobenzene	EPA 8260B	6K22012	0.48	1.0	ND	1	11/22/06	11/22/06	
1,1,2-Trichloroethane	EPA 8260B	6K22012	0.30	1.0	ND	1	11/22/06	11/22/06	
1,1,1-Trichloroethane	EPA 8260B	6K22012	0.30	1.0	ND	1	11/22/06	11/22/06	
Trichlorofluoromethane	EPA 8260B	6K22012	0.34	2.0	ND	1	11/22/06	11/22/06	
1,2,3-Trichloropropane	EPA 8260B	6K22012	0.40	1.0	ND	1	11/22/06	11/22/06	
1,2,4-Trimethylbenzene	EPA 8260B	6K22012	0.23	1.0	ND	1	11/22/06	11/22/06	
1,3,5-Trimethylbenzene	EPA 8260B	6K22012	0.26	1.0	ND	1	11/22/06	11/22/06	
Vinyl acetate	EPA 8260B	6K22012	1.7	6.0	ND	1	11/22/06	11/22/06	
Vinyl chloride	EPA 8260B	6K22012	0.30	0.50	ND	1	11/22/06	11/22/06	
Xylenes, Total	EPA 8260B	6K22012	0.90	1.0	ND	1	11/22/06	11/22/06	
Surrogate: 4-Bromofluorobenzene (80-120%)					94 %				
Surrogate: Dibromofluoromethane (80-120%)					97 %				
Surrogate: Toluene-d8 (80-120%)					97 %				

TestAmerica - Irvine, CA
Nicholas Marz For Michele Chamberlin
Project Manager

11/24/07

The results pertain only to the samples tested in the laboratory. This report shall not be reproduced, except in full, without written permission from TestAmerica.

IPK2310 <Page 16 of 44>

TestAmerica

ANALYTICAL TESTING CORPORATION

TAIT Environmental/Boeing 701 N. Parkcenter Drive Santa Ana, CA 92705 Attention: Mehmet Pehlivan	Project ID: Boeing C-6 Torrance EM2727 Report Number: IPK2310	Sampled: 11/20/06 Received: 11/20/06
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VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IPK2310-06RE1 (MWC016_WG112006_0001 - Water) - cont.									
Reporting Units: ug/l									
Trichloroethene	EPA 8260B	6K22012	2.6	10	1100	10	11/22/06	11/22/06	
Surrogate: 4-Bromofluorobenzene (80-120%)					93 %				
Surrogate: Dibromofluoromethane (80-120%)					100 %				
Surrogate: Toluene-d8 (80-120%)					95 %				

TestAmerica - Irvine, CA
 Nicholas Marz For Michele Chamberlin
 Project Manager

Mof 2/07

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IPK2310 <Page 17 of 44>

LDC #: 16470B1
 SDG #: IPK2310
 Laboratory: Test America

VALIDATION COMPLETENESS WORKSHEET
 EPA Region 1 - Tier 1

Date: 3/20/07
 Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: GC/MS Volatiles (EPA SW 846 Method 8260B)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	Δ	Sampling dates: 11/20/04
II.	GC/MS Instrument performance check	N	
III.	Initial calibration	N	
IV.	Continuing calibration	N	
V.	Blanks	SW	
VI.	Surrogate spikes	A	
VII.	Matrix spike/Matrix spike duplicates	SW	
VIII.	Laboratory control samples	A	LC9
IX.	Regional Quality Assurance and Quality Control	N	
X.	Internal standards	N	
XI.	Target compound identification	N	
XII.	Compound quantitation/CRQLs	N	
XIII.	Tentatively identified compounds (TICs)	N	
XIV.	System performance	N	
XV.	Overall assessment of data	A	
XVI.	Field duplicates	NSW	D = 1 + 3 2 + 4
XVII.	Field blanks	N	

Note: A = Acceptable
 N = Not provided/applicable
 SW = See worksheet
 ND = No compounds detected
 R = Rinsate
 FB = Field blank
 D = Duplicate
 TB = Trip blank
 EB = Equipment blank

Validated Samples: *water*

1	MWC015_WG112006_0001	11	CR22012	21	31
2	MWC015_WG112006_0001RE1	12		22	32
3	MWC016_WG112006_0001	13		23	33
4	MWC016_WG112006_0001RE1	14		24	34
5	MWC015_WG112006_0001MS	15		25	35
6	MWC015_WG112006_0001MSD	16		26	36
7		17		27	37
8		18		28	38
9		19		29	39
10		20		30	40

TARGET COMPOUND WORKSHEET

METHOD: VOA (EPA SW 846 Method 8260B)

A. Chloromethane*	S. Trichloroethene	KK. Trichlorofluoromethane	CCC. tert-Butylbenzene	UUU. 1,2-Dichlorotetrafluoroethane
B. Bromomethane	T. Dibromochloromethane	LL. Methyl-tert-butyl ether	DDD. 1,2,4-Trimethylbenzene	VVV. 4-Ethyltoluene
C. Vinyl chloride**	U. 1,1,2-Trichloroethane	MM. 1,2-Dibromo-3-chloropropane	EEE. sec-Butylbenzene	WWW. Ethanol
D. Chloroethane	V. Benzene	NN. Methyl ethyl ketone	FFF. 1,3-Dichlorobenzene	XXX. Di-isopropyl ether
E. Methylene chloride	W. trans-1,3-Dichloropropene	OO. 2,2-Dichloropropane	GGG. p-Isopropyltoluene	YYY. tert-Butanol
F. Acetone	X. Bromoform*	PP. Bromochloromethane	HHH. 1,4-Dichlorobenzene	ZZZ. tert-Butyl alcohol
G. Carbon disulfide	Y. 4-Methyl-2-pentanone	QQ. 1,1-Dichloropropene	III. n-Butylbenzene	AAAA. Ethyl tert-butyl ether
H. 1,1-Dichloroethane**	Z. 2-Hexanone	RR. Dibromomethane	JJJ. 1,2-Dichlorobenzene	BBBB. tert-Amyl methyl ether
I. 1,1-Dichloroethane*	AA. Tetrachloroethene	SS. 1,3-Dichloropropane	KKK. 1,2,4-Trichlorobenzene	CCCC. 1-Chlorohexane
J. 1,2-Dichloroethene, total	BB. 1,1,2,2-Tetrachloroethane*	TT. 1,2-Dibromoethane	LLL. Hexachlorobutadiene	DDDD. Isopropyl alcohol
K. Chloroform*	CC. Toluene**	UU. 1,1,1,2-Tetrachloroethane	MMM. Naphthalene	EEEE. Acetonitrile
L. 1,2-Dichloroethane	DD. Chlorobenzene*	VV. Isopropylbenzene	NNN. 1,2,3-Trichlorobenzene	FFFF. Acrolein
M. 2-Butanone	EE. Ethylbenzene**	WW. Bromobenzene	OOO. 1,3,5-Trichlorobenzene	GGGG. Acrylonitrile
N. 1,1,1-Trichloroethane	FF. Styrene	XX. 1,2,3-Trichloropropane	PPP. trans-1,2-Dichloroethene	HHHH. 1,4-Dioxane
O. Carbon tetrachloride	GG. Xylenes, total	YY. n-Propylbenzene	QQQ. cis-1,2-Dichloroethene	IIII. Isobutyl alcohol
P. Bromodichloromethane	HH. Vinyl acetate	ZZ. 2-Chlorotoluene	RRR. m,p-Xylenes	JJJJ. Methacrylonitrile
Q. 1,2-Dichloropropane**	II. 2-Chloroethylvinyl ether	AAA. 1,3,5-Trimethylbenzene	SSS. o-Xylene	KKKK. Propionitrile
R. cis-1,3-Dichloropropane	JJ. Dichlorodifluoromethane	BBB. 4-Chlorotoluene	TTT. 1,1,2-Trichloro-1,2,2-trifluoroethane	LLLL.

* = System performance check compounds (SPCC) for RRF ; ** = Calibration check compounds (CCC) for %RSD.

LDC #: 16470B1
 SDG #: 1PK2310

VALIDATION FINDINGS WORKSHEET
Blanks

Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: GC/MS VOA (EPA SW 846 Method 8260B)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

X N N/A Was a method blank associated with every sample in this SDG?
Y N N/A Was a method blank analyzed at least once every 12 hours for each matrix and concentration?
Y N N/A Was there contamination in the method blanks? If yes, please see the qualifications below.

Blank analysis date: 11/22/06

Conc. units: ug/L Associated Samples: All (MP)

Compound	Blank ID	Sample Identification
Tetrahydrofuran	6K2292-B/K	
Methylene chloride	4.41	
Acetone		
CRQL		

Blank analysis date: _____
 Conc. units: _____

Associated Samples: _____

Compound	Blank ID	Sample Identification
Methylene chloride		
Acetone		
CRQL		

All results were qualified using the criteria stated below except those circled.

Note: Common contaminants such as Methylene chloride, Acetone, 2-Butanone, Carbon disulfide and TlCs that were detected in samples within ten times the associated method blank concentration were qualified as not detected, "U". Other contaminants within five times the method blank concentration were also qualified as not detected, "U".

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Boeing Realty Corp. C-6, EM2727
Collection Date: November 21, 2006
LDC Report Date: March 28, 2007
Matrix: Water
Parameters: Dissolved Manganese
Validation Level: Tier 2 & 3
Laboratory: TestAmerica

Sample Delivery Group (SDG): IPK2470

Sample Identification

IWC001_WG112106_0001**
MWC024_WG112106_001

**Indicates sample underwent Tier 3 review

Introduction

This data review covers 2 water samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 6010B for Dissolved Manganese.

This review follows a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Inorganic Data Review (October 2004) as there are no current guidelines for the methods stated above.

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

Blanks are summarized in Section III.

Field duplicates are summarized in Section XIII.

Samples indicated by a double asterisk on the front cover underwent a Tier 3 review. A Tier 2 review was performed on all of the other samples. Raw data were not evaluated for the samples reviewed by Tier 2 criteria since this review is based on QC data.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- N Presumptive evidence of presence of the constituent.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. Calibration

An initial calibration was performed.

The frequency and analysis criteria of the initial calibration verification (ICV) and continuing calibration verification (CCV) were met.

III. Blanks

Method blanks were reviewed for each matrix as applicable. No contaminant concentrations were found in the initial, continuing and preparation blanks.

IV. ICP Interference Check Sample (ICS) Analysis

The frequency of analysis was met.

The criteria for analysis were met.

V. Matrix Spike Analysis

Matrix spike (MS) and matrix spike duplicate (MSD) samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

VI. Duplicate Sample Analysis

Duplicate (DUP) sample analyses were reviewed for each matrix as applicable.

VII. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) were within QC limits.

VIII. Internal Standards

ICP-MS was not utilized in this SDG.

IX. Furnace Atomic Absorption QC

Graphite furnace atomic absorption was not utilized in this SDG.

X. ICP Serial Dilution

ICP serial dilution was not performed for this SDG.

XI. Sample Result Verification

All sample result verifications were acceptable for samples on which a Tier 3 review was performed. Raw data were not evaluated for the samples reviewed by Tier 2 criteria.

XII. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

XIII. Field Duplicates

No field duplicates were identified in this SDG.

XIV. Field Blanks

No field blanks were identified in this SDG.

PLP

**Boeing Realty Corp. C-6, EM2727
Dissolved Manganese - Data Qualification Summary - SDG IPK2470**

No Sample Data Qualified in this SDG

**Boeing Realty Corp. C-6, EM2727
Dissolved Manganese - Laboratory Blank Data Qualification Summary - SDG
IPK2470**

No Sample Data Qualified in this SDG

TestAmerica

ANALYTICAL TESTING CORPORATION

17461 Derian Avenue, Suite 100, Irvine, CA 92614 (949) 261-1022 Fax: (949) 260-3297

TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727

Report Number: IPK2470

Sampled: 11/21/06
Received: 11/21/06

DISSOLVED METALS

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IPK2470-07 (IWC001_WG112106_0001 - Water)									
Reporting Units: mg/l									
Manganese	EPA 6010B-Diss	6K21150	0.0070	0.020	0.038	1	11/21/06	11/22/06	
Sample ID: IPK2470-09 (MWC024_WG112106_0001 - Water)									
Reporting Units: mg/l									
Manganese	EPA 6010B-Diss	6K21150	0.0070	0.020	0.015	1	11/21/06	11/22/06	J

TestAmerica - Irvine, CA
Michele Chamberlin
Project Manager

11/20/07

The results pertain only to the samples tested in the laboratory. This report shall not be reproduced, except in full, without written permission from TestAmerica.

IPK2470 <Page 34 of 62>

LDC #: 16470A4
 SDG #: IPK2470
 Laboratory: Test America

VALIDATION COMPLETENESS WORKSHEET
 EPA Region 1 - Tier 2/3

Date: 3/27/07
 Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: Dissolved Manganese (EPA SW 846 Method 6010B)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: 11/21/06
II.	Calibration	A	
III.	Blanks	A	
IV.	ICP Interference Check Sample (ICS) Analysis	A	
V.	Matrix Spike Analysis	A	none detected
VI.	Duplicate Sample Analysis	N	
VII.	Laboratory Control Samples (LCS)	A	LCS
VIII.	Internal Standard (ICP-MS)	N	not utilized
IX.	Furnace Atomic Absorption QC	N	
X.	ICP Serial Dilution	N	not performed
XI.	Sample Result Verification	A	Not reviewed for Tier II validation.
XII.	Overall Assessment of Data	A	
XIII.	Field Duplicates	N	
XIV.	Field Blanks	N	

Note: A = Acceptable ND = No compounds detected D = Duplicate
 N = Not provided/applicable R = Rinsate TB = Trip blank
 SW = See worksheet FB = Field blank EB = Equipment blank

Validated Samples: ** Indicates sample underwent Tier III validation

1	IWC001_WG112106_0001**	11		21		31	
2	MWC024_WG112106_001	12		22		32	
3	PS	13		23		33	
4		14		24		34	
5		15		25		35	
6		16		26		36	
7		17		27		37	
8		18		28		38	
9		19		29		39	
10		20		30		40	

Notes: _____

LDC #: 1647-AY
 SDG #: See cover

VALIDATION FINDINGS CHECKLIST

Page: 1 of 2
 Reviewer: HW
 2nd Reviewer: A

Method: Metals (EPA SW 846 Method 6010/7000/6020)

Validation Area	Yes	No	NA	Findings/Comments
I: Technical Holding Times				
All technical holding times were met.	✓			
Cooler temperature criteria was met.	✓			
II: Calibration				
Were all instruments calibrated daily, each set-up time?	✓			
Were the proper number of standards used?	✓			
Were all initial and continuing calibration verification %Rs within the 90-110% (80-120% for mercury and 85-115% for cyanide) QC limits?	✓			
Were all initial calibration correlation coefficients > 0.995? (Level IV only)	✓			
III: Blanks				
Was a method blank associated with every sample in this SDG?	✓			
Was there contamination in the method blanks? If yes, please see the Blanks validation completeness worksheet.		✓		
IV: Interference Check Sample				
Were ICP interference check samples performed daily?	✓			
Were the AB solution percent recoveries (%R) with the 80-120% QC limits?	✓			
V: Matrix Spike/Matrix Spike Duplicates				
Were a matrix spike (MS) and duplicate (DUP) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD or MS/DUP. Soil / Water.	✓			
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the 75-125 QC limits? If the sample concentration exceeded the spike concentration by a factor of 4 or more, no action was taken.	✓			
Were the MS/MSD or duplicate relative percent differences (RPD) ≤ 20% for waters and ≤ 35% for soil samples? A control limit of +/- RL (+/- 2X RL for soil) was used for samples that were < 5X the RL, including when only one of the duplicate sample values were < 5X the RL.	✓			
VI: Laboratory Control Samples				
Was an LCS analyzed for this SDG?	✓			
Was an LCS analyzed per extraction batch?	✓			
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the 80-120% QC limits for water samples and laboratory established QC limits for soils?	✓			
VII: For use: Atomic Absorption QC				
If MSA was performed, was the correlation coefficients > 0.995?			✓	
Do all applicable analyses have duplicate injections? (Level IV only)			✓	
For sample concentrations > RL, are applicable duplicate injection RSD values < 20%? (Level IV only)			✓	
Were analytical spike recoveries within the 85-115% QC limits?			✓	

LDC #: 16470A4
 SDG #: See call

VALIDATION FINDINGS CHECKLIST

Page: 2 of 2
 Reviewer: LM
 2nd Reviewer: [Signature]

Validation Area	Yes	No	NA	Findings/Comments
VI. ICP Serial Dilutions				
Was an ICP serial dilution analyzed if analyte concentrations were > 50X the IDL?		✓		
Were all percent differences (%Ds) < 10%?			✓	
Was there evidence of negative interference? If yes, professional judgement will be used to qualify the data.			✓	
VII. Internal Standards (EPA SW-846 Method 6020)				
Were all the percent recoveries (%R) within the 30-120% of the intensity of the internal standard in the associated initial calibration?			✓	
If the %Rs were outside the criteria, was a reanalysis performed?			✓	
VIII. Regional Quality Assurance and Quality Control				
Were performance evaluation (PE) samples performed?			✓	
Were the performance evaluation (PE) samples within the acceptance limits?			✓	
IX. Sample Basis Verification				
Were RLs adjusted to reflect all sample dilutions and dry weight factors applicable to level IV validation?	✓			
X. Overall Assessment of data				
Overall assessment of data was found to be acceptable.	✓			
XI. Field Duplicates				
Field duplicate pairs were identified in this SDG.		✓		
Target analytes were detected in the field duplicates.			✓	
XII. Field blanks				
Field blanks were identified in this SDG.		✓		
Target analytes were detected in the field blanks.			✓	

LDC #: 1647-AF
 SDG #: See case

VALIDATION FINDINGS WORKSHEET
Initial and Continuing Calibration Calculation Verification

Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: Trace Metals (EPA SW 846 Method 6010/7000)

An initial and continuing calibration verification percent recovery (%R) was recalculated for each type of analysis using the following formula:

$\%R = \frac{\text{Found}}{\text{True}} \times 100$ Where: Found = concentration (in ug/L) of each analyte measured in the analysis of the ICV or CCV solution
 True = concentration (in ug/L) of each analyte in the ICV or CCV source

Standard ID	Type of Analysis	Element	Found (ug/L)	True (ug/L)	Recalculated		Reported		Acceptable (Y/N)
					%R	%R	%R	%R	
ICV	ICP (Initial calibration)	Mn	1.986	2.00	99		MR		Y
	GFAA (Initial calibration)								
	CVAA (Initial calibration)								
CCV	ICP (Continuing calibration)	Mn	1.01	1.00	101		MR		Y
	GFAA (Continuing calibration)								
	CVAA (Continuing calibration)								
	Cyanide (Initial calibration)								
	Cyanide (Continuing calibration)								

Comments: Refer to Calibration Verification findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Boeing Realty Corp. C-6, EM2727
Collection Date: November 21, 2006
LDC Report Date: April 5, 2007
Matrix: Water
Parameters: Volatiles
Validation Level: Tier 2 & 3
Laboratory: TestAmerica, Inc.

Sample Delivery Group (SDG): IPK2470

Sample Identification

IWC002_WG112106_0001
IWC002_WG112106_0001RE1
IWC002_WG112106_0001RE2
IWC001_WG112106_0001**
IWC001_WG112106_0001RE1**
IWC001_WG112106_0001RE2**
MWC024_WG112106_001
MWC024_WG112106_001RE1
MWC024_WG112106_001RE2
IWC002_WG112106_0001MS
IWC002_WG112106_0001MSD

**Indicates sample underwent Tier 3 review

Introduction

This data review covers 11 water samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 8260B for Volatiles.

This review follows a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Organic Data Review (October 1999) as there are no current guidelines for the method stated above.

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

Blank results are summarized in Section V.

Field duplicates are summarized in Section XVI.

Samples indicated by a double asterisk on the front cover underwent a Tier Level 3 review. A Tier 2 review was performed on all of the other samples. Raw data were not evaluated for the samples reviewed by Tier 2 criteria since this review is based on QC data.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- N Presumptive evidence of presence of the constituent.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. GC/MS Instrument Performance Check

Instrument performance was checked at 12 hour intervals.

All ion abundance requirements were met.

III. Initial Calibration

Initial calibration was performed using required standard concentrations.

Percent relative standard deviations (%RSD) were less than or equal to 15.0% for each individual compound and less than or equal to 30.0% for calibration check compounds (CCCs).

In the case where %RSD was greater than 15.0%, the laboratory used a calibration curve to evaluate the compound. All coefficients of determination (r^2) were greater than or equal to 0.990 .

For the purposes of technical evaluation, all compounds were evaluated against the 30.0% (%RSD) National Functional Guideline criteria. Unless noted above, all compounds were within the validation criteria.

Average relative response factors (RRF) for all volatile target compounds and system performance check compounds (SPCCs) were within method and validation criteria.

IV. Continuing Calibration

Continuing calibration was performed at the required frequencies.

Percent differences (%D) between the initial calibration RRF and the continuing calibration RRF were within the method criteria of less than or equal to 20.0% for calibration check compounds (CCCs).

For the purposes of technical evaluation, all compounds were evaluated against the 25.0% (%D) National Functional Guideline criteria. Unless noted above, all compounds were within the validation criteria with the following exceptions:

Date	Compound	%D	Associated Samples	Flag	A or P
11/22/06 (20:10)	Tetrahydrofuran	48	MWC024_WG112106_001 6K22027-BLK1	J (all detects) UJ (all non-detects)	A

All of the continuing calibration RRF values were within method and validation criteria with the following exceptions:

Date	Compound	RRF (Limits)	Associated Samples	Flag	A or P
11/22/06 (09:39)	2-Butanone	0.048 (≥0.05)	IWC002_WG112106_0001 IWC001_WG112106_0001** IWC002_WG112106_0001MS IWC002_WG112106_0001MSD 6K22017-BLK1	J (all detects) UJ (all non-detects)	A

V. Blanks

Method blanks were reviewed for each matrix as applicable. No volatile contaminants were found in the method blanks with the following exceptions:

Method Blank ID	Analysis Date	Compound TIC (RT in minutes)	Concentration	Associated Samples
6K22017-BLK1	11/22/06	Tetrahydrofuran	4.83 ug/L	IWC002_WG112106_0001 IWC001_WG112106_0001**
6K22027-BLK1	11/22/06	Tetrahydrofuran	5.50 ug/L	MWC024_WG112106_001

Sample concentrations were compared to concentrations detected in the method blanks. The sample concentrations were either not detected or were significantly greater (>10X for common contaminants, >5X for other contaminants) than the concentrations found in the associated method blanks.

VI. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VII. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

VIII. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) were within QC limits.

IX. Regional Quality Assurance and Quality Control

Not applicable.

X. Internal Standards

All internal standard areas and retention times were within QC limits.

XI. Target Compound Identifications

All target compound identifications were within validation criteria for samples on which a Tier 3 review was performed. Raw data were not evaluated for the samples reviewed by Tier 2 criteria.

XII. Compound Quantitation and CRQLs

All compound quantitation and CRQLs were within validation criteria for samples on which a Tier 3 review was performed. Raw data were not evaluated for the samples reviewed by Tier 2 criteria.

XIII. Tentatively Identified Compounds (TICs)

Tentatively identified compounds were not reported by the laboratory.

XIV. System Performance

The system performance was acceptable for samples on which a Tier 3 review was performed. Raw data were not evaluated for the samples reviewed by Tier 2 criteria.

XV. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

XVI. Field Duplicates

No field duplicates were identified in this SDG.

XVII. Field Blanks

No field blanks were identified in this SDG.

Boeing Realty Corp. C-6, EM2727
Volatiles - Data Qualification Summary - SDG IPK2470

SDG	Sample	Compound	Flag	A or P	Reason
IPK2470	MWC024_WG112106_001	Tetrahydrofuran	J (all detects) UJ (all non-detects)	A	Continuing calibration (%D)
IPK2470	IWC002_WG112106_0001 IWC001_WG112106_0001**	2-Butanone	J (all detects) UJ (all non-detects)	A	Continuing calibration (RRF)

Boeing Realty Corp. C-6, EM2727
Volatiles - Laboratory Blank Data Qualification Summary - SDG IPK2470

No Sample Data Qualified in this SDG

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ANALYTICAL TESTING CORPORATION

17461 Derian Avenue, Suite 100, Irvine, CA 92614 (949) 261-1022 Fax: (949) 260-3297

TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IPK2470

Sampled: 11/21/06
Received: 11/21/06

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IPK2470-05 (FWC002_WG112106_0001 - Water)									
Reporting Units: ug/l									
Acetone	EPA 8260B	6K22017	4.5	10	ND	1	11/22/06	11/22/06	
Benzene	EPA 8260B	6K22017	0.28	1.0	0.30	1	11/22/06	11/22/06	J
Bromobenzene	EPA 8260B	6K22017	0.27	1.0	ND	1	11/22/06	11/22/06	
Bromochloromethane	EPA 8260B	6K22017	0.32	1.0	ND	1	11/22/06	11/22/06	
Bromodichloromethane	EPA 8260B	6K22017	0.30	1.0	ND	1	11/22/06	11/22/06	
Bromoform	EPA 8260B	6K22017	0.40	1.0	ND	1	11/22/06	11/22/06	
Bromomethane	EPA 8260B	6K22017	0.42	1.0	ND	1	11/22/06	11/22/06	
2-Butanone (MEK)	EPA 8260B	6K22017	3.8	5.0	ND	1	11/22/06	11/22/06	
n-Butylbenzene	EPA 8260B	6K22017	0.37	1.0	ND	1	11/22/06	11/22/06	
sec-Butylbenzene	EPA 8260B	6K22017	0.25	1.0	ND	1	11/22/06	11/22/06	
tert-Butylbenzene	EPA 8260B	6K22017	0.22	1.0	ND	1	11/22/06	11/22/06	
Carbon Disulfide	EPA 8260B	6K22017	0.48	1.0	ND	1	11/22/06	11/22/06	
Carbon tetrachloride	EPA 8260B	6K22017	0.28	0.50	ND	1	11/22/06	11/22/06	
Chlorobenzene	EPA 8260B	6K22017	0.36	1.0	ND	1	11/22/06	11/22/06	
Chloroethane	EPA 8260B	6K22017	0.40	2.0	ND	1	11/22/06	11/22/06	
Chloroform	EPA 8260B	6K22017	0.33	1.0	21	1	11/22/06	11/22/06	
Chloromethane	EPA 8260B	6K22017	0.40	2.0	ND	1	11/22/06	11/22/06	
2-Chlorotoluene	EPA 8260B	6K22017	0.28	1.0	ND	1	11/22/06	11/22/06	
4-Chlorotoluene	EPA 8260B	6K22017	0.29	1.0	ND	1	11/22/06	11/22/06	
1,2-Dibromo-3-chloropropane	EPA 8260B	6K22017	0.97	2.0	ND	1	11/22/06	11/22/06	
Dibromochloromethane	EPA 8260B	6K22017	0.28	1.0	ND	1	11/22/06	11/22/06	
1,2-Dibromoethane (EDB)	EPA 8260B	6K22017	0.40	1.0	ND	1	11/22/06	11/22/06	
1,4-Dichlorobenzene	EPA 8260B	6K22017	0.37	1.0	ND	1	11/22/06	11/22/06	
1,2-Dichlorobenzene	EPA 8260B	6K22017	0.32	1.0	ND	1	11/22/06	11/22/06	
1,3-Dichlorobenzene	EPA 8260B	6K22017	0.35	1.0	ND	1	11/22/06	11/22/06	
Dichlorodifluoromethane	EPA 8260B	6K22017	0.79	1.0	ND	1	11/22/06	11/22/06	
1,2-Dichloroethane	EPA 8260B	6K22017	0.28	0.50	3.9	1	11/22/06	11/22/06	
1,1-Dichloroethane	EPA 8260B	6K22017	0.27	1.0	5.9	1	11/22/06	11/22/06	
1,1-Dichloroethene	EPA 8260B	6K22017	0.42	1.0	110	1	11/22/06	11/22/06	
cis-1,2-Dichloroethene	EPA 8260B	6K22017	0.32	1.0	9.2	1	11/22/06	11/22/06	
trans-1,2-Dichloroethene	EPA 8260B	6K22017	0.27	1.0	4.1	1	11/22/06	11/22/06	
1,2-Dichloropropane	EPA 8260B	6K22017	0.35	1.0	ND	1	11/22/06	11/22/06	
2,2-Dichloropropane	EPA 8260B	6K22017	0.34	1.0	ND	1	11/22/06	11/22/06	
cis-1,3-Dichloropropene	EPA 8260B	6K22017	0.22	0.50	ND	1	11/22/06	11/22/06	
1,1-Dichloropropene	EPA 8260B	6K22017	0.28	1.0	ND	1	11/22/06	11/22/06	
trans-1,3-Dichloropropene	EPA 8260B	6K22017	0.32	0.50	ND	1	11/22/06	11/22/06	
Ethylbenzene	EPA 8260B	6K22017	0.25	1.0	ND	1	11/22/06	11/22/06	
Hexachlorobutadiene	EPA 8260B	6K22017	0.38	1.0	ND	1	11/22/06	11/22/06	
2-Hexanone	EPA 8260B	6K22017	2.6	6.0	ND	1	11/22/06	11/22/06	
Iodomethane	EPA 8260B	6K22017	1.0	2.0	ND	1	11/22/06	11/22/06	
Isopropylbenzene	EPA 8260B	6K22017	0.25	1.0	ND	1	11/22/06	11/22/06	

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Lichele Chamberlin
Project Manager

11/24/07

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TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IPK2470

Sampled: 11/21/06
Received: 11/21/06

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IPK2470-05 (IWC002_WG112106_0001 - Water) - cont.									
Reporting Units: ug/l									
p-Isopropyltoluene	EPA 8260B	6K22017	0.28	1.0	ND	1	11/22/06	11/22/06	
Methyl-tert-butyl Ether (MTBE)	EPA 8260B	6K22017	0.32	1.0	ND	1	11/22/06	11/22/06	
Methylene chloride	EPA 8260B	6K22017	0.95	1.0	ND	1	11/22/06	11/22/06	
4-Methyl-2-pentanone (MIBK)	EPA 8260B	6K22017	3.5	5.0	ND	1	11/22/06	11/22/06	
n-Propylbenzene	EPA 8260B	6K22017	0.27	1.0	ND	1	11/22/06	11/22/06	
Styrene	EPA 8260B	6K22017	0.16	1.0	ND	1	11/22/06	11/22/06	
1,1,1,2-Tetrachloroethane	EPA 8260B	6K22017	0.27	1.0	ND	1	11/22/06	11/22/06	
1,1,2,2-Tetrachloroethane	EPA 8260B	6K22017	0.24	1.0	ND	1	11/22/06	11/22/06	
Tetrachloroethene	EPA 8260B	6K22017	0.32	1.0	0.98	1	11/22/06	11/22/06	J
Tetrahydrofuran (THF)	EPA 8260B	6K22017	3.5	10	ND	1	11/22/06	11/22/06	
Toluene	EPA 8260B	6K22017	0.36	1.0	ND	1	11/22/06	11/22/06	
1,2,3-Trichlorobenzene	EPA 8260B	6K22017	0.30	1.0	ND	1	11/22/06	11/22/06	
1,2,4-Trichlorobenzene	EPA 8260B	6K22017	0.48	1.0	ND	1	11/22/06	11/22/06	
1,1,2-Trichloroethane	EPA 8260B	6K22017	0.30	1.0	2.4	1	11/22/06	11/22/06	
1,1,1-Trichloroethane	EPA 8260B	6K22017	0.30	1.0	ND	1	11/22/06	11/22/06	
Trichlorofluoromethane	EPA 8260B	6K22017	0.34	2.0	ND	1	11/22/06	11/22/06	
1,2,3-Trichloropropane	EPA 8260B	6K22017	0.40	1.0	ND	1	11/22/06	11/22/06	
1,2,4-Trimethylbenzene	EPA 8260B	6K22017	0.23	1.0	ND	1	11/22/06	11/22/06	
1,3,5-Trimethylbenzene	EPA 8260B	6K22017	0.26	1.0	ND	1	11/22/06	11/22/06	
Vinyl chloride	EPA 8260B	6K22017	0.30	0.50	ND	1	11/22/06	11/22/06	
Xylenes, Total	EPA 8260B	6K22017	0.90	1.0	ND	1	11/22/06	11/22/06	
Surrogate: 4-Bromofluorobenzene (80-120%)					102 %				
Surrogate: Dibromofluoromethane (80-120%)					108 %				
Surrogate: Toluene-d8 (80-120%)					100 %				

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Michele Chamberlin
Project Manager

11/21/07

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TAIT Environmental/Boeing 701 N. Parkcenter Drive Santa Ana, CA 92705 Attention: Mehmet Pehlivan	Project ID: Boeing C-6 Torrance EM2727 Report Number: IPK2470	Sampled: 11/21/06 Received: 11/21/06
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VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IPK2470-05RE1 (IWC002_WG112106_0001 - Water) - cont.									
Reporting Units: ug/l									
Trichloroethene	EPA 8260B	6K22031	5.2	20	3000	20	11/22/06	11/23/06	
Surrogate: 4-Bromofluorobenzene (80-120%)					82 %				
Surrogate: Dibromofluoromethane (80-120%)					99 %				
Surrogate: Toluene-d8 (80-120%)					100 %				

TestAmerica - Irvine, CA
Sichele Chamberlin
Project Manager

M. Pehlivan

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ANALYTICAL TESTING CORPORATION

17461 Derian Avenue, Suite 100, Irvine, CA 92614 (949) 261-1022 Fax: (949) 260-3297

TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IPK2470

Sampled: 11/21/06
Received: 11/21/06

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IPK2470-05RE2 (TWC002_WG112106_0001 - Water) - cont.									
Reporting Units: ug/l									
Vinyl acetate	EPA 8260B	6K27030	1.7	6.0	ND	1	11/27/06	11/27/06	
Surrogate: 4-Bromofluorobenzene (80-120%)					91 %				
Surrogate: Dibromofluoromethane (80-120%)					89 %				
Surrogate: Toluene-d8 (80-120%)					95 %				

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Michele Chamberlin
Project Manager

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IPK2470 <Page 17 of 62>

TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IPK2470

Sampled: 11/21/06
Received: 11/21/06

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IPK2470-07 (IWC001_WG112106_0001 - Water)									
Reporting Units: ug/l									
Acetone	EPA 8260B	6K22017	4.5	10	ND	1	11/22/06	11/22/06	
Benzene	EPA 8260B	6K22017	0.28	1.0	0.43	1	11/22/06	11/22/06	J
Bromobenzene	EPA 8260B	6K22017	0.27	1.0	ND	1	11/22/06	11/22/06	
Bromochloromethane	EPA 8260B	6K22017	0.32	1.0	ND	1	11/22/06	11/22/06	
Bromodichloromethane	EPA 8260B	6K22017	0.30	1.0	ND	1	11/22/06	11/22/06	
Bromoform	EPA 8260B	6K22017	0.40	1.0	ND	1	11/22/06	11/22/06	
Bromomethane	EPA 8260B	6K22017	0.42	1.0	ND	1	11/22/06	11/22/06	
2-Butanone (MEK)	EPA 8260B	6K22017	3.8	5.0	ND	1	11/22/06	11/22/06	
n-Butylbenzene	EPA 8260B	6K22017	0.37	1.0	ND	1	11/22/06	11/22/06	
sec-Butylbenzene	EPA 8260B	6K22017	0.25	1.0	ND	1	11/22/06	11/22/06	
tert-Butylbenzene	EPA 8260B	6K22017	0.22	1.0	ND	1	11/22/06	11/22/06	
Carbon Disulfide	EPA 8260B	6K22017	0.48	1.0	ND	1	11/22/06	11/22/06	
Carbon tetrachloride	EPA 8260B	6K22017	0.28	0.50	0.97	1	11/22/06	11/22/06	
Chlorobenzene	EPA 8260B	6K22017	0.36	1.0	ND	1	11/22/06	11/22/06	
Chloroethane	EPA 8260B	6K22017	0.40	2.0	ND	1	11/22/06	11/22/06	
Chloromethane	EPA 8260B	6K22017	0.40	2.0	ND	1	11/22/06	11/22/06	
2-Chlorotoluene	EPA 8260B	6K22017	0.28	1.0	ND	1	11/22/06	11/22/06	
4-Chlorotoluene	EPA 8260B	6K22017	0.29	1.0	ND	1	11/22/06	11/22/06	
1,2-Dibromo-3-chloropropane	EPA 8260B	6K22017	0.97	2.0	ND	1	11/22/06	11/22/06	
Dibromochloromethane	EPA 8260B	6K22017	0.28	1.0	ND	1	11/22/06	11/22/06	
1,2-Dibromoethane (EDB)	EPA 8260B	6K22017	0.40	1.0	ND	1	11/22/06	11/22/06	
1,4-Dichlorobenzene	EPA 8260B	6K22017	0.37	1.0	ND	1	11/22/06	11/22/06	
1,2-Dichlorobenzene	EPA 8260B	6K22017	0.32	1.0	ND	1	11/22/06	11/22/06	
1,3-Dichlorobenzene	EPA 8260B	6K22017	0.35	1.0	ND	1	11/22/06	11/22/06	
Dichlorodifluoromethane	EPA 8260B	6K22017	0.79	1.0	ND	1	11/22/06	11/22/06	
1,2-Dichloroethane	EPA 8260B	6K22017	0.28	0.50	1.2	1	11/22/06	11/22/06	
1,1-Dichloroethane	EPA 8260B	6K22017	0.27	1.0	1.1	1	11/22/06	11/22/06	
1,1-Dichloroethene	EPA 8260B	6K22017	0.42	1.0	53	1	11/22/06	11/22/06	
cis-1,2-Dichloroethene	EPA 8260B	6K22017	0.32	1.0	1.6	1	11/22/06	11/22/06	
trans-1,2-Dichloroethene	EPA 8260B	6K22017	0.27	1.0	0.33	1	11/22/06	11/22/06	J
1,2-Dichloropropane	EPA 8260B	6K22017	0.35	1.0	ND	1	11/22/06	11/22/06	
2,2-Dichloropropane	EPA 8260B	6K22017	0.34	1.0	ND	1	11/22/06	11/22/06	
cis-1,3-Dichloropropene	EPA 8260B	6K22017	0.22	0.50	ND	1	11/22/06	11/22/06	
1,1-Dichloropropene	EPA 8260B	6K22017	0.28	1.0	ND	1	11/22/06	11/22/06	
trans-1,3-Dichloropropene	EPA 8260B	6K22017	0.32	0.50	ND	1	11/22/06	11/22/06	
Ethylbenzene	EPA 8260B	6K22017	0.25	1.0	ND	1	11/22/06	11/22/06	
Hexachlorobutadiene	EPA 8260B	6K22017	0.38	1.0	ND	1	11/22/06	11/22/06	
2-Hexanone	EPA 8260B	6K22017	2.6	6.0	ND	1	11/22/06	11/22/06	
Iodomethane	EPA 8260B	6K22017	1.0	2.0	ND	1	11/22/06	11/22/06	
Isopropylbenzene	EPA 8260B	6K22017	0.25	1.0	ND	1	11/22/06	11/22/06	
p-Isopropyltoluene	EPA 8260B	6K22017	0.28	1.0	ND	1	11/22/06	11/22/06	

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Fichele Chamberlin
Project Manager

11/21/06

TAIT Environmental/Boeing 701 N. Parkcenter Drive Santa Ana, CA 92705 Attention: Mehmet Pehlivan	Project ID: Boeing C-6 Torrance EM2727 Report Number: IPK2470	Sampled: 11/21/06 Received: 11/21/06
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VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IPK2470-07 (IWC001_WG112106_0001 - Water) - cont.									
Reporting Units: ug/l									
Methyl-tert-butyl Ether (MTBE)	EPA 8260B	6K22017	0.32	1.0	ND	1	11/22/06	11/22/06	
Methylene chloride	EPA 8260B	6K22017	0.95	1.0	ND	1	11/22/06	11/22/06	
4-Methyl-2-pentanone (MIBK)	EPA 8260B	6K22017	3.5	5.0	ND	1	11/22/06	11/22/06	
n-Propylbenzene	EPA 8260B	6K22017	0.27	1.0	ND	1	11/22/06	11/22/06	
Styrene	EPA 8260B	6K22017	0.16	1.0	ND	1	11/22/06	11/22/06	
1,1,1,2-Tetrachloroethane	EPA 8260B	6K22017	0.27	1.0	ND	1	11/22/06	11/22/06	
1,1,2,2-Tetrachloroethane	EPA 8260B	6K22017	0.24	1.0	ND	1	11/22/06	11/22/06	
Tetrachloroethene	EPA 8260B	6K22017	0.32	1.0	2.6	1	11/22/06	11/22/06	
Tetrahydrofuran (THF)	EPA 8260B	6K22017	3.5	10	ND	1	11/22/06	11/22/06	
Toluene	EPA 8260B	6K22017	0.36	1.0	ND	1	11/22/06	11/22/06	
1,2,3-Trichlorobenzene	EPA 8260B	6K22017	0.30	1.0	ND	1	11/22/06	11/22/06	
1,2,4-Trichlorobenzene	EPA 8260B	6K22017	0.48	1.0	ND	1	11/22/06	11/22/06	
1,1,2-Trichloroethane	EPA 8260B	6K22017	0.30	1.0	0.59	1	11/22/06	11/22/06	J
1,1,1-Trichloroethane	EPA 8260B	6K22017	0.30	1.0	0.33	1	11/22/06	11/22/06	J
Trichlorofluoromethane	EPA 8260B	6K22017	0.34	2.0	ND	1	11/22/06	11/22/06	
1,2,3-Trichloropropane	EPA 8260B	6K22017	0.40	1.0	ND	1	11/22/06	11/22/06	
1,2,4-Trimethylbenzene	EPA 8260B	6K22017	0.23	1.0	ND	1	11/22/06	11/22/06	
1,3,5-Trimethylbenzene	EPA 8260B	6K22017	0.26	1.0	ND	1	11/22/06	11/22/06	
Vinyl chloride	EPA 8260B	6K22017	0.30	0.50	ND	1	11/22/06	11/22/06	
Xylenes, Total	EPA 8260B	6K22017	0.90	1.0	ND	1	11/22/06	11/22/06	
Surrogate: 4-Bromofluorobenzene (80-120%)					102 %				
Surrogate: Dibromofluoromethane (80-120%)					110 %				
Surrogate: Toluene-d8 (80-120%)					101 %				

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Michele Chamberlin
Project Manager

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TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IPK2470

Sampled: 11/21/06
Received: 11/21/06

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IPK2470-07RE1 (IWC001_WG112106_0001 - Water) - cont.									
Reporting Units: ug/l									
Chloroform	EPA 8260B	6K22031	6.6	20	730	20	11/22/06	11/23/06	
Trichloroethene	EPA 8260B	6K22031	5.2	20	2600	20	11/22/06	11/23/06	
Surrogate: 4-Bromofluorobenzene (80-120%)					82 %				
Surrogate: Dibromofluoromethane (80-120%)					98 %				
Surrogate: Toluene-d8 (80-120%)					101 %				

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Project Manager

11/21/06

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TAIT Environmental/Boeing
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Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IPK2470

Sampled: 11/21/06
Received: 11/21/06

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IPK2470-07RE2 (IWC001_WG112106_0001 - Water) - cont.									
Reporting Units: ug/l									
Vinyl acetate	EPA 8260B	6K27030	1.7	6.0	ND	1	11/27/06	11/27/06	
Surrogate: 4-Bromofluorobenzene (80-120%)					92 %				
Surrogate: Dibromofluoromethane (80-120%)					89 %				
Surrogate: Toluene-d8 (80-120%)					97 %				

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TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IPK2470

Sampled: 11/21/06
Received: 11/21/06

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IPK2470-09 (MWC024_WG112106_0001 - Water)									
Reporting Units: ug/l									
Acetone	EPA 8260B	6K22027	4.5	10	ND	1	11/22/06	11/22/06	
Benzene	EPA 8260B	6K22027	0.28	1.0	0.66	1	11/22/06	11/22/06	J
Bromobenzene	EPA 8260B	6K22027	0.27	1.0	ND	1	11/22/06	11/22/06	
Bromochloromethane	EPA 8260B	6K22027	0.32	1.0	ND	1	11/22/06	11/22/06	
Bromodichloromethane	EPA 8260B	6K22027	0.30	1.0	ND	1	11/22/06	11/22/06	
Bromoform	EPA 8260B	6K22027	0.40	1.0	ND	1	11/22/06	11/22/06	
Bromomethane	EPA 8260B	6K22027	0.42	1.0	ND	1	11/22/06	11/22/06	
2-Butanone (MEK)	EPA 8260B	6K22027	3.8	5.0	ND	1	11/22/06	11/22/06	
n-Butylbenzene	EPA 8260B	6K22027	0.37	1.0	ND	1	11/22/06	11/22/06	
sec-Butylbenzene	EPA 8260B	6K22027	0.25	1.0	ND	1	11/22/06	11/22/06	
tert-Butylbenzene	EPA 8260B	6K22027	0.22	1.0	ND	1	11/22/06	11/22/06	
Carbon Disulfide	EPA 8260B	6K22027	0.48	1.0	ND	1	11/22/06	11/22/06	
Carbon tetrachloride	EPA 8260B	6K22027	0.28	0.50	0.94	1	11/22/06	11/22/06	
Chlorobenzene	EPA 8260B	6K22027	0.36	1.0	0.53	1	11/22/06	11/22/06	J
Chloroethane	EPA 8260B	6K22027	0.40	2.0	ND	1	11/22/06	11/22/06	
Chloromethane	EPA 8260B	6K22027	0.40	2.0	ND	1	11/22/06	11/22/06	
2-Chlorotoluene	EPA 8260B	6K22027	0.28	1.0	ND	1	11/22/06	11/22/06	
4-Chlorotoluene	EPA 8260B	6K22027	0.29	1.0	ND	1	11/22/06	11/22/06	
1,2-Dibromo-3-chloropropane	EPA 8260B	6K22027	0.97	2.0	ND	1	11/22/06	11/22/06	
Dibromochloromethane	EPA 8260B	6K22027	0.28	1.0	ND	1	11/22/06	11/22/06	
1,2-Dibromoethane (EDB)	EPA 8260B	6K22027	0.40	1.0	ND	1	11/22/06	11/22/06	
1,4-Dichlorobenzene	EPA 8260B	6K22027	0.37	1.0	ND	1	11/22/06	11/22/06	
1,2-Dichlorobenzene	EPA 8260B	6K22027	0.32	1.0	ND	1	11/22/06	11/22/06	
1,3-Dichlorobenzene	EPA 8260B	6K22027	0.35	1.0	ND	1	11/22/06	11/22/06	
Dichlorodifluoromethane	EPA 8260B	6K22027	0.79	1.0	ND	1	11/22/06	11/22/06	
1,2-Dichloroethane	EPA 8260B	6K22027	0.28	0.50	0.65	1	11/22/06	11/22/06	
1,1-Dichloroethane	EPA 8260B	6K22027	0.27	1.0	1.1	1	11/22/06	11/22/06	
1,1-Dichloroethene	EPA 8260B	6K22027	0.42	1.0	33	1	11/22/06	11/22/06	
cis-1,2-Dichloroethene	EPA 8260B	6K22027	0.32	1.0	1.9	1	11/22/06	11/22/06	
trans-1,2-Dichloroethene	EPA 8260B	6K22027	0.27	1.0	0.57	1	11/22/06	11/22/06	J
1,2-Dichloropropane	EPA 8260B	6K22027	0.35	1.0	ND	1	11/22/06	11/22/06	
2,2-Dichloropropane	EPA 8260B	6K22027	0.34	1.0	ND	1	11/22/06	11/22/06	
cis-1,3-Dichloropropene	EPA 8260B	6K22027	0.22	0.50	ND	1	11/22/06	11/22/06	
1,1-Dichloropropene	EPA 8260B	6K22027	0.28	1.0	ND	1	11/22/06	11/22/06	
trans-1,3-Dichloropropene	EPA 8260B	6K22027	0.32	0.50	ND	1	11/22/06	11/22/06	
Ethylbenzene	EPA 8260B	6K22027	0.25	1.0	ND	1	11/22/06	11/22/06	
Hexachlorobutadiene	EPA 8260B	6K22027	0.38	1.0	ND	1	11/22/06	11/22/06	
2-Hexanone	EPA 8260B	6K22027	2.6	6.0	ND	1	11/22/06	11/22/06	
Iodomethane	EPA 8260B	6K22027	1.0	2.0	ND	1	11/22/06	11/22/06	
Isopropylbenzene	EPA 8260B	6K22027	0.25	1.0	ND	1	11/22/06	11/22/06	
p-Isopropyltoluene	EPA 8260B	6K22027	0.28	1.0	ND	1	11/22/06	11/22/06	

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Michele Chamberlin
Project Manager

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ANALYTICAL TESTING CORPORATION

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TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IPK2470

Sampled: 11/21/06
Received: 11/21/06

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IPK2470-09 (MWC024_WG112106_0001 - Water) - cont.									
Reporting Units: ug/l									
Methyl-tert-butyl Ether (MTBE)	EPA 8260B	6K22027	0.32	1.0	ND	1	11/22/06	11/22/06	
Methylene chloride	EPA 8260B	6K22027	0.95	1.0	ND	1	11/22/06	11/22/06	
4-Methyl-2-pentanone (MIBK)	EPA 8260B	6K22027	3.5	5.0	ND	1	11/22/06	11/22/06	
n-Propylbenzene	EPA 8260B	6K22027	0.27	1.0	ND	1	11/22/06	11/22/06	
Styrene	EPA 8260B	6K22027	0.16	1.0	ND	1	11/22/06	11/22/06	
1,1,1,2-Tetrachloroethane	EPA 8260B	6K22027	0.27	1.0	ND	1	11/22/06	11/22/06	
1,1,2,2-Tetrachloroethane	EPA 8260B	6K22027	0.24	1.0	ND	1	11/22/06	11/22/06	
Tetrachloroethene	EPA 8260B	6K22027	0.32	1.0	1.2	1	11/22/06	11/22/06	
Tetrahydrofuran (THF)	EPA 8260B	6K22027	3.5	10	ND	1	11/22/06	11/22/06	
Toluene	EPA 8260B	6K22027	0.36	1.0	ND	1	11/22/06	11/22/06	
1,2,3-Trichlorobenzene	EPA 8260B	6K22027	0.30	1.0	ND	1	11/22/06	11/22/06	
1,2,4-Trichlorobenzene	EPA 8260B	6K22027	0.48	1.0	ND	1	11/22/06	11/22/06	
1,1,2-Trichloroethane	EPA 8260B	6K22027	0.30	1.0	0.44	1	11/22/06	11/22/06	J
1,1,1-Trichloroethane	EPA 8260B	6K22027	0.30	1.0	ND	1	11/22/06	11/22/06	
Trichlorofluoromethane	EPA 8260B	6K22027	0.34	2.0	ND	1	11/22/06	11/22/06	
1,2,3-Trichloropropane	EPA 8260B	6K22027	0.40	1.0	ND	1	11/22/06	11/22/06	
1,2,4-Trimethylbenzene	EPA 8260B	6K22027	0.23	1.0	ND	1	11/22/06	11/22/06	
1,3,5-Trimethylbenzene	EPA 8260B	6K22027	0.26	1.0	ND	1	11/22/06	11/22/06	
Vinyl chloride	EPA 8260B	6K22027	0.30	0.50	ND	1	11/22/06	11/22/06	
Xylenes, Total	EPA 8260B	6K22027	0.90	1.0	ND	1	11/22/06	11/22/06	
Surrogate: 4-Bromofluorobenzene (80-120%)					101 %				
Surrogate: Dibromofluoromethane (80-120%)					110 %				
Surrogate: Toluene-d8 (80-120%)					100 %				

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Michele Chamberlin
Project Manager

11/20/07

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TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IPK2470

Sampled: 11/21/06
Received: 11/21/06

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IPK2470-09RE1 (MWC024_WG112106_0001 - Water) - cont.									
Reporting Units: ug/l									
Chloroform	EPA 8260B	6K25014	6.6	20	460	20	11/25/06	11/25/06	
Trichloroethene	EPA 8260B	6K25014	5.2	20	2800	20	11/25/06	11/25/06	
Surrogate: 4-Bromofluorobenzene (80-120%)					106 %				
Surrogate: Dibromofluoromethane (80-120%)					114 %				
Surrogate: Toluene-d8 (80-120%)					109 %				

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Nicole Chamberlin
Project Manager

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IPK2470 <Page 32 of 62>

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Project ID: Boeing C-6 Torrance
EM2727
Report Number: IPK2470

Sampled: 11/21/06
Received: 11/21/06

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IPK2470-09RE2 (MWC024_WG112106_0001 - Water) - cont.									
Reporting Units: ug/l									
Vinyl acetate	EPA 8260B	6K24024	1.7	6.0	ND	1	11/24/06	11/24/06	
Surrogate: 4-Bromofluorobenzene (80-120%)					92 %				
Surrogate: Dibromofluoromethane (80-120%)					95 %				
Surrogate: Toluene-d8 (80-120%)					97 %				

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Michele Chamberlin
Project Manager

11/24/06

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IPK2470 <Page 33 of 62>

LDC #: 16470A1
 SDG #: IPK2470
 Laboratory: Test America

VALIDATION COMPLETENESS WORKSHEET
 EPA Region 1 - Tier 2/3

Date: 3/3/07
 Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: GC/MS Volatiles (EPA SW 846 Method 8260B)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: 11/21/06
II.	GC/MS Instrument performance check	A	
III.	Initial calibration	A	% RSD, r ² 20-990
IV.	Continuing calibration	SW	
V.	Blanks	SW	
VI.	Surrogate spikes	A	
VII.	Matrix spike/Matrix spike duplicates	A	
VIII.	Laboratory control samples	A	LCS
IX.	Regional Quality Assurance and Quality Control	N	
X.	Internal standards	A	
XI.	Target compound identification	A	Not reviewed for Tier II validation.
XII.	Compound quantitation/CRQLs	A	Not reviewed for Tier II validation.
XIII.	Tentatively identified compounds (TICs)	N	Not reviewed for Tier II validation. not reported
XIV.	System performance	A	Not reviewed for Tier II validation.
XV.	Overall assessment of data	A	
XVI.	Field duplicates	N	SW D = 1, 2, 3, 4, 5, 6
XVII.	Field blanks	N	

Note: A = Acceptable
 N = Not provided/applicable
 SW = See worksheet
 *ND = No compounds detected
 R = Rinsate
 FB = Field blank
 D = Duplicate
 TB = Trip blank
 EB = Equipment blank

Validated Samples: **Indicates sample underwent Tier III validation

Water

1	IWC002_WG112106_0001	11	IWC002_WG112106_0001MSD	21	6K22017 X	31
2	IWC002_WG112106_0001RE1	12		22	6K22031 X	32
3	IWC002_WG112106_0001RE2	13		23	6K27030 X	33
4	IWC001_WG112106_0001**	14		24	6K22027 X	34
5	IWC001_WG112106_0001RE1**	15		25	6K25014 X	35
6	IWC001_WG112106_0001RE2**	16		26	6K24024 X	36
7	MWC024_WG112106_001	17		27		37
8	MWC024_WG112106_001RE1	18		28		38
9	MWC024_WG112106_001RE2	19		29		39
10	IWC002_WG112106_0001MS	20		30		40

DC #: 16470-A1
 SDG #: see cover

VALIDATION FINDINGS CHECKLIST

Page: 1 of 2
 Reviewer: PJ
 2nd Reviewer: KL

Method: Volatiles (EPA SW 846 Method 8260B)

Validation Area	Yes	No	NA	Findings/Comments
All technical holding times were met.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Cooler temperature criteria was met.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the BFB performance results reviewed and found to be within the specified criteria?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all samples analyzed within the 12 hour clock criteria?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Did the laboratory perform a 5 point calibration prior to sample analysis?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all percent relative standard deviations (%RSD) and relative response factors (RRF) within method criteria for all CCCs and SPCCs?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a curve fit used for evaluation?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Did the initial calibration meet the curve fit acceptance criteria of > 0.990?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all percent relative standard deviations (%RSD) ≤ 30% and relative response factors (RRF) > 0.05?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a continuing calibration standard analyzed at least once every 12 hours for each instrument?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all percent differences (%D) and relative response factors (RRF) within method criteria for all CCCs and SPCCs?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all percent differences (%D) ≤ 25% and relative response factors (RRF) ≥ 0.05?	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
Was a method blank associated with every sample in this SDG?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a method blank analyzed at least once every 12 hours for each matrix and concentration?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was there contamination in the method blanks? If yes, please see the Blanks validation completeness worksheet.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all surrogate %R within QC limits?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
If the percent recovery (%R) for one or more surrogates was out of QC limits, was a reanalysis performed to confirm samples with %R outside of criteria?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were a matrix spike (MS) and matrix spike duplicate (MSD) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD. Soil / Water.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a MS/MSD analyzed every 20 samples of each matrix?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the QC limits?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was an LCS analyzed for this SDG?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	

DC #: 16470A
 IG #: see cover

VALIDATION FINDINGS CHECKLIST

Page: 2 of 2
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

Validation Area	Yes	No	NA	Findings/Comments
Was an LCS analyzed per analytical batch?	/			
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the QC limits?	/			
Performance Evaluation (PE) Samples				
Were performance evaluation (PE) samples performed?			/	
Were the performance evaluation (PE) samples within the acceptance limits?			/	
Internal Standard Area Counts				
Were internal standard area counts within -50% or +100% of the associated calibration standard?	/			
Were retention times within + 30 seconds of the associated calibration standard?	/			
Retention Times				
Were relative retention times (RRT's) within + 0.06 RRT units of the standard?	/			
Did compound spectra meet specified EPA "Functional Guidelines" criteria?	/			
Were chromatogram peaks verified and accounted for?	/			
Internal Standard (IS) Quantitation				
Were the correct internal standard (IS), quantitation ion and relative response factor (RRF) used to quantitate the compound?	/			
Were compound quantitation and CRQLs adjusted to reflect all sample dilutions and dry weight factors applicable to level IV validation?	/			
Reference Spectra				
Were the major ions (> 10 percent relative intensity) in the reference spectrum evaluated in sample spectrum?			/	
Were relative intensities of the major ions within ± 20% between the sample and the reference spectra?			/	
Did the raw data indicate that the laboratory performed a library search for all required peaks in the chromatograms (samples and blanks)?			/	
System Performance				
System performance was found to be acceptable.	/			
Overall Assessment of Data				
Overall assessment of data was found to be acceptable.	/			
Field Duplicates				
Field duplicate pairs were identified in this SDG.		/		
Target compounds were detected in the field duplicates.			/	
Field Blanks				
Field blanks were identified in this SDG.		/		
Target compounds were detected in the field blanks.			/	

TARGET COMPOUND WORKSHEET

METHOD: VOA (EPA SW 846 Method 8260B)

A. Chloromethane*	S. Trichloroethane	KK. Trichlorofluoromethane	CCC. tert-Butylbenzene	UUU. 1,2-Dichlorotetrafluoroethane
B. Bromomethane	T. Dibromochloromethane	LL. Methyl-tert-butyl ether	DDD. 1,2,4-Trimethylbenzene	VVV. 4-Ethyltoluene
C. Vinyl chloride**	U. 1,1,2-Trichloroethane	MM. 1,2-Dibromo-3-chloropropane	EEE. sec-Butylbenzene	WWW. Ethanol
D. Chloroethane	V. Benzene	NN. Methyl ethyl ketone	FFF. 1,3-Dichlorobenzene	XXX. Di-isopropyl ether
E. Methylene chloride	W. trans-1,3-Dichloropropene	OO. 2,2-Dichloropropane	GGG. p-Isopropyltoluene	YYY. tert-Butanol
F. Acetone	X. Bromoform*	PP. Bromochloromethane	HHH. 1,4-Dichlorobenzene	ZZZ. tert-Butyl alcohol
G. Carbon disulfide	Y. 4-Methyl-2-pentanone	QQ. 1,1-Dichloropropene	III. n-Butylbenzene	AAA. Ethyl tert-butyl ether
H. 1,1-Dichloroethane**	Z. 2-Hexanone	RR. Dibromomethane	JJJ. 1,2-Dichlorobenzene	BBB. tert-Amyl methyl ether
I. 1,1-Dichloroethane*	AA. Tetrachloroethane	SS. 1,3-Dichloropropane	KKK. 1,2,4-Trichlorobenzene	CCC. 1-Chlorohexane
J. 1,2-Dichloroethane, total	BB. 1,1,2,2-Tetrachloroethane*	TT. 1,2-Dibromoethane	LLL. Hexachlorobutadiene	DDD. Isopropyl alcohol
K. Chloroform**	CC. Toluene**	UU. 1,1,1,2-Tetrachloroethane	MMM. Naphthalene	EEE. Acetonitrile
L. 1,2-Dichloroethane	DD. Chlorobenzene*	VV. Isopropylbenzene	NNN. 1,2,3-Trichlorobenzene	FFF. Acrolein
M. 2-Butanone	EE. Ethylbenzene**	WW. Bromobenzene	OOO. 1,3,5-Trichlorobenzene	GGG. Acrylonitrile
N. 1,1,1-Trichloroethane	FF. Styrene	XX. 1,2,3-Trichloropropane	PPP. trans-1,2-Dichloroethane	HHH. 1,4-Dioxane
O. Carbon tetrachloride	GG. Xylenes, total	YY. n-Propylbenzene	QQQ. cis-1,2-Dichloroethane	III. Isobutyl alcohol
P. Bromodichloromethane	HH. Vinyl acetate	ZZ. 2-Chlorotoluene	RRR. m,p-Xylenes	JJJ. Methacrylonitrile
Q. 1,2-Dichloropropane**	II. 2-Chloroethylvinyl ether	AAA. 1,3,5-Trimethylbenzene	SSS. o-Xylene	KKK. Propionitrile
R. cis-1,3-Dichloropropene	JJ. Dichlorodifluoromethane	BBB. 4-Chlorotoluene	TTT. 1,1,2-Trichloro-1,2,2-trifluoroethane	LLL. LLL.

* = System performance check compounds (SPCC) for RRF; ** = Calibration check compounds (CCC) for %RSD.

LDC #: 10470A

SDG #: for each

VALIDATION FINDINGS WORKSHEET

Initial Calibration Calculation Verification

Page: 1 of 1
Reviewer: [Signature]
2nd Reviewer: [Signature]

METHOD: GC/MS VOA (EPA SW 846 Method 8260B)

The Relative Response Factor (RRF), average RRF, and percent relative standard deviation (%RSD) were recalculated for the compounds identified below using the following calculations:

$RRF = (A_s/C_s)/(A_i/C_i)$
 average RRF = sum of the RRFs/number of standards
 $\%RSD = 100 * (S/X)$

A_s = Area of compound,
 C_s = Concentration of compound,
 S = Standard deviation of the RRFs
 X = Mean of the RRFs

A_i = Area of associated internal standard
 C_i = Concentration of internal standard

#	Standard ID	Calibration Date	Compound (Reference/Internal Standard)	Reported		Recalculated		Reported		Recalculated	
				RRF (25 std)	RRF (25 std)	RRF (25 std)	RRF (25 std)	Average RRF (Initial)	%RSD	Average RRF (Initial)	%RSD
1	GCMS33	11/15/06	Methylene chloride (1st internal standard)	0.455	0.455	0.455	0.455	0.442	5.53	0.442	5.53
			Trichlorethene (2nd internal standard)	0.410	0.410	0.410	0.410	0.385	9.22	0.385	9.22
			1,2-DCE Toluene (3rd internal standard)	1.429	1.429	1.429	1.429	1.379	9.63	1.379	9.63
2	GCMS58	11/15/06	Methylene chloride (1st internal standard)	1.418	1.418	1.418	1.418	1.337	10.92	1.337	10.92
			Trichlorethene (2nd internal standard)								
			Toluene (3rd internal standard)								
3	GCMS56	11/15/06	TCE	0.386	0.386	0.386	0.386	0.365	5.81	0.365	5.81
			Methylene chloride (1st internal standard)	0.992	0.992	0.992	0.992	0.977	5.86	0.977	5.86
			Chloroform Trichloroethene (2nd internal standard)								
			Toluene (3rd internal standard)								
4	GCMS55	11/23/06	Methylene chloride (1st internal standard)	0.798	0.798	0.798	0.798	0.679	14.85	0.679	14.85
			Vinyl Acetate Trichlorethene (2nd internal standard)								
			Toluene (3rd internal standard)								

Comments: Refer to Initial Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

INCLC.15B

LDC #: 164170A
 SDG #: for control

VALIDATION FINDINGS WORKSHEET
Continuing Calibration Results Verification

Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: GC/MS VOA (EPA SW 846 Method 8260B)

The percent difference (%D) of the initial calibration average Relative Response Factors (RRFs) and the continuing calibration RRFs were recalculated for the compounds identified below using the following calculation:

% Difference = $100 * (\text{ave. RRF} - \text{RRF}) / \text{ave. RRF}$ Where: ave. RRF = initial calibration average RRF
 RRF = $(A_s / C_s) / (A_i / C_i)$ RRF = continuing calibration RRF
 A_s = Area of compound, A_i = Area of associated internal standard
 C_s = Concentration of compound, C_i = Concentration of internal standard

#	Standard ID	Calibration Date	Compound (Reference Internal Standard)	Average RRF (Initial)	Reported		Recalculated	
					RRF (CC)	%D	RRF (CC)	%D
1	CW 9:39 AM (33)	11/22/07	Methylene chloride (1st internal standard)	0.442	0.400	9.5	0.400	9.5
			Trichloroethene (2nd internal standard)	0.385	0.391	1.6	0.391	1.6
			Toluene (3rd internal standard) Ethyl Benzene	1.379	1.410	2.2	1.410	2.2
2			Methylene chloride (1st internal standard) 1,2-DCB	1.337	1.403	4.9	1.403	4.9
			Trichloroethene (2nd internal standard)					
			Toluene (3rd internal standard)					
3	CW 6:48 PM (52)	11/22/07	Methylene chloride (1st internal standard) CWs refer to Methylene chloride (1st internal standard)	0.977	1.014	4.0	1.016	4.0
			Trichloroethene (2nd internal standard)	0.365	0.358	1.9	0.358	1.9
			Toluene (3rd internal standard)					
4	CW 4:41 PM (55)	11/27/07	Methylene chloride (1st internal standard) Methylene chloride (1st internal standard)	0.679	0.682	0.4	0.682	0.4
			Trichloroethene (2nd internal standard)					
			Toluene (3rd internal standard)					

Comments: Refer to Continuing Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

CONCL:15B

LDC #: 16470A1
 SDG #: μ cones

VALIDATION FINDINGS WORKSHEET
Surrogate Results Verification

Page: 1 of 1
 Reviewer: [Signature]
 2nd reviewer: [Signature]

METHOD: GC/MS VOA (EPA SW 846 Method 8260B)

The percent recoveries (%R) of surrogates were recalculated for the compounds identified below using the following calculation:

% Recovery: SF/SS * 100

Where: SF = Surrogate Found
 SS = Surrogate Spiked

Sample ID: 4

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Toluene-d8	25.0	25.31	101	101	0
Bromofluorobenzene	↓	25.62	102	102	↓
1,2-Dichloroethane-d4					
Dibromofluoromethane	↓	27.40	110	110	↓

Sample ID: _____

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Toluene-d8					
Bromofluorobenzene					
1,2-Dichloroethane-d4					
Dibromofluoromethane					

Sample ID: _____

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Toluene-d8					
Bromofluorobenzene					
1,2-Dichloroethane-d4					
Dibromofluoromethane					

Sample ID: _____

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Toluene-d8					
Bromofluorobenzene					
1,2-Dichloroethane-d4					
Dibromofluoromethane					

Sample ID: _____

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Toluene-d8					
Bromofluorobenzene					
1,2-Dichloroethane-d4					
Dibromofluoromethane					

LDC #: 16470A
 SDG #: per cover

VALIDATION FINDINGS WORKSHEET
Matrix Spike/Matrix Spike Duplicates Results Verification

Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: GC/MS VOA (EPA SW 846 Method 8260B)

The percent recoveries (%R) and Relative Percent Difference (RPD) of the matrix spike and matrix spike duplicate were recalculated for the compounds identified below using the following calculation:

% Recovery = $100 \cdot (SSC - SC) / SA$

Where: SSC = Spiked sample concentration
 SA = Spike added

SC = Sample concentration

RPD = $100 \cdot MSC - MSDC / (MSC + MSDC)$

MSC = Matrix spike percent recovery

MSDC = Matrix spike duplicate percent recovery

MS/MSD sample: 10 + 11

Compound	Spike Added (ug/l)		Sample Concentration (ug/l)	Spiked Sample Concentration (ug/l)		Matrix Spike Percent Recovery		Matrix Spike Duplicate Percent Recovery		MS/MSD RPD	
	MS	MSD		MS	MSD	Reported	Recalc.	Reported	Recalc.	Reported	Recalculated
1,1-Dichloroethene	26.0	26.0	11.0	13.0	12.6	80	80	64	64	3	3
Trichloroethene	NA		NA	NA							
Benzene			0.30	24.4	24.8	96	96	98	98	2	2
Toluene			ND	24.8	24.9	99	99	100	100	0	0
Chlorobenzene			ND	25.7	25.9	103	103	104	104	1	1

Comments: Refer to Matrix Spike/Matrix Spike Duplicates findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Boeing Realty Corp. C-6, EM2727
Collection Date: November 21, 2006
LDC Report Date: March 28, 2007
Matrix: Water
Parameters: Wet Chemistry
Validation Level: Tier 2 & 3
Laboratory: TestAmerica

Sample Delivery Group (SDG): IPK2470

Sample Identification

IWC001_WG112106_0001**
MWC024_WG112106_001

**Indicates sample underwent Tier 3 review

Introduction

This data review covers 2 water samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA Method 300.0 for Bromide, Chloride, Nitrate, Nitrite, and Sulfate, EPA Method 310.1 for Alkalinity, and EPA Method 415.1 Total Organic Carbon.

The review follows a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Inorganic Data Review (October 2004) as there are no current guidelines for the methods stated above.

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

Blank results are summarized in Section III.

Field duplicates are summarized in Section IX.

Samples indicated by a double asterisk on the front cover underwent a Tier 3 review. A Tier 2 review was performed on all of the other samples. Raw data were not evaluated for the samples reviewed by Tier 2 criteria since this review is based on QC data.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- N Presumptive evidence of presence of the constituent.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. Calibration

a. Initial Calibration

All criteria for the initial calibration of each method were met.

b. Calibration Verification

Calibration verification frequency and analysis criteria were met for each method when applicable.

III. Blanks

Method blanks were reviewed for each matrix as applicable. No contaminant concentrations were found in the initial, continuing and preparation blanks with the following exceptions:

Method Blank ID	Analyte	Concentration	Associated Samples
MB	Sulfate Chloride	0.165 mg/L 0.220 mg/L	All samples in SDG 1PK2470
ICB/CCB	Chloride	0.194 mg/L 0.189 mg/L	All samples in SDG 1PK2470

Sample concentrations were compared to concentrations detected in the method blanks. The sample concentrations were either not detected or were significantly greater (>5X blank contaminants) than the concentrations found in the associated method blanks.

IV. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

V. Duplicates

Duplicate (DUP) sample analyses were reviewed for each matrix as applicable. Results were within QC limits.

VI. Laboratory Control Samples

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) were within QC limits.

VII. Sample Result Verification

All sample result verifications were acceptable for samples on which a Tier 3 review was performed. Raw data were not evaluated for the samples reviewed by Tier 2 criteria.

VIII. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

IX. Field Duplicates

No field duplicates were identified in this SDG.

X. Field Blanks

No field blanks were identified in this SDG.

**Boeing Realty Corp. C-6, EM2727
Wet Chemistry - Data Qualification Summary - SDG IPK2470**

No Sample Data Qualified in this SDG

**Boeing Realty Corp. C-6, EM2727
Wet Chemistry - Laboratory Blank Data Qualification Summary - SDG IPK2470**

No Sample Data Qualified in this SDG

TestAmerica

ANALYTICAL TESTING CORPORATION

17461 Derian Avenue, Suite 100, Irvine, CA 92614 (949) 261-1022 Fax: (949) 260-3297

TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IPK2470

Sampled: 11/21/06
Received: 11/21/06

INORGANICS

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IPK2470-07 (IWC001_WG112106_0001 - Water)									
Reporting Units: mg/l									
Alkalinity as CaCO3	EPA 310.1	6L04068	2.0	2.0	200	1	12/04/06	12/04/06	
Bromide	EPA 300.0	6L01049	1.8	2.5	2.8	5	12/01/06	12/01/06	
Chloride	EPA 300.0	6L01049	5.0	25	280	50	12/01/06	12/02/06	
Nitrate-NO3	EPA 300.0	6K22067	0.25	0.50	17	1	11/22/06	11/22/06	
Nitrite-NO2	EPA 300.0	6K22067	0.30	0.50	ND	1	11/22/06	11/22/06	
Sulfate	EPA 300.0	6K22067	0.15	0.50	52	1	11/22/06	11/22/06	
Total Organic Carbon	EPA 415.1	6L03024	0.50	1.0	ND	1	12/03/06	12/03/06	
Sample ID: IPK2470-09 (MWC024_WG112106_0001 - Water)									
Reporting Units: mg/l									
Alkalinity as CaCO3	EPA 310.1	6L04068	2.0	2.0	180	1	12/04/06	12/04/06	
Bromide	EPA 300.0	6L01049	1.8	2.5	2.6	5	12/01/06	12/02/06	
Chloride	EPA 300.0	6L01049	5.0	25	480	50	12/01/06	12/02/06	
Nitrate-NO3	EPA 300.0	6K22067	0.25	0.50	25	1	11/22/06	11/22/06	
Nitrite-NO2	EPA 300.0	6K22067	0.30	0.50	ND	1	11/22/06	11/22/06	
Sulfate	EPA 300.0	6K22067	0.15	0.50	37	1	11/22/06	11/22/06	
Total Organic Carbon	EPA 415.1	6L03024	0.50	1.0	ND	1	12/03/06	12/03/06	

TestAmerica - Irvine, CA
Michele Chamberlin
Project Manager

The results pertain only to the samples tested in the laboratory. This report shall not be reproduced, except in full, without written permission from TestAmerica.

IPK2470 <Page 35 of 62>

LDC #: 16470A6
 SDG #: IPK2470
 Laboratory: Test America

VALIDATION COMPLETENESS WORKSHEET
 EPA Region 1 - Tier 2/3

Date: 3/27/07
 Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: Alkalinity (EPA Method 310.1), Bromide, Chloride, Nitrate ~~N~~, Nitrite ~~N~~, Sulfate (EPA Method 300.0),
 TOC (EPA Method 415.1)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: 4/21/06
IIa.	Initial calibration	A	
IIb.	Calibration verification	A	
III.	Blanks	SW	
IVa.	Matrix Spike/(Matrix Spike) Duplicates	A	MS/MSD/OSP
IVb.	Laboratory control samples	A	LCS
V.	Sample result verification	A	Not reviewed for Tier II validation.
VI.	Overall assessment of data	A	
VII.	Field duplicates	N	
VIII.	Field blanks	N	

Note: A = Acceptable ND = No compounds detected D = Duplicate
 N = Not provided/applicable R = Rinsate TB = Trip blank
 SW = See worksheet FB = Field blank EB = Equipment blank

Validated Samples: ** Indicates sample underwent Tier III validation

1	IWC001_WG112106_0001**	11		21		31	
2	MWC024_WG112106_001	12		22		32	
3	MB	13		23		33	
4		14		24		34	
5		15		25		35	
6		16		26		36	
7		17		27		37	
8		18		28		38	
9		19		29		39	
10		20		30		40	

Notes: _____

LDC #: 16490A6
 SDG #: 7PK2470

VALIDATION FINDINGS CHECKLIST

Page: 1 of 1
 Reviewer: MLY
 2nd Reviewer: R

Method: Inorganics (EPA Method See cover)

Validation Area	Yes	No	NA	Findings/Comments
II. Technical Holding Times				
All technical holding times were met.	✓			
Cooler temperature criteria was met.	✓			
III. Calibration				
Were all instruments calibrated daily, each set-up time?	✓			
Were the proper number of standards used?	✓			
Were all initial calibration correlation coefficients > 0.995?	✓			
Were all initial and continuing calibration verification %Rs within the 90-110% QC limits?	✓			
Were titrant checks performed as required? (Level IV only)			✓	
Were balance checks performed as required? (Level IV only)	✓			
IV. Blanks				
Was a method blank associated with every sample in this SDG?	✓			
Was there contamination in the method blanks? If yes, please see the Blanks validation completeness worksheet.	✓			
V. Matrix Spike, Matrix and Duplicate, and Duplicate				
Were a matrix spike (MS) and duplicate (DUP) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD or MS/DUP. Soil / Water.	✓			
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the 75-125 QC limits? If the sample concentration exceeded the spike concentration by a factor of 4 or more, no action was taken.	✓			
Were the MS/MSD or duplicate relative percent differences (RPD) ≤ 20% for waters and ≤ 35% for soil samples? A control limit of ≤ CRDL (≤ 2X CRDL for soil) was used for samples that were ≤ 5X the CRDL, including when only one of the duplicate sample values were ≤ 5X the CRDL.	✓			
VI. Laboratory Controls and QC				
Was an LCS analyzed for this SDG?	✓			
Was an LCS analyzed per extraction batch?	✓			
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the 80-120% (85-115% for Method 300.0) QC limits?	✓			
VII. Regional Quality Assurance and Quality Control				
Were performance evaluation (PE) samples performed?			✓	
Were the performance evaluation (PE) samples within the acceptance limits?			✓	

LDC #: 16490A6
 SDG #: See cover

VALIDATION FINDINGS CHECKLIST

Page: 1 of 1
 Reviewer: WY
 2nd Reviewer: [Signature]

Validation Area	Yes	No	NA	Findings/Comments
Wf Sample Result Verification				
Were RLs adjusted to reflect all sample dilutions and dry weight factors applicable to level IV validation?	✓			
Were detection limits < RL?	✓			
Overall Assessment				
Overall assessment of data was found to be acceptable.	✓			
Field Duplicate Pairs				
Field duplicate pairs were identified in this SDG.		✓		
Target analytes were detected in the field duplicates.			✓	
Field Blanks				
Field blanks were identified in this SDG.		✓		
Target analytes were detected in the field blanks.			✓	

LDC #: 1647046
 SDG #: See cover

VALIDATION FINDINGS WORKSHEET
Sample Specific Analysis Reference

Page: 1 of 1
 Reviewer: MM
 2nd reviewer: A

All circled methods are applicable to each sample.

Sample ID	Parameter
<u>112</u>	pH TDS <u>(C)</u> F <u>(C)</u> NO ₃ <u>(C)</u> NO ₂ <u>(C)</u> SO ₄ PO ₄ ALK CN NH ₃ TKN <u>(C)</u> TOC CR ⁰⁺ <u>(B)</u>
	pH TDS CI F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS CI F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS CI F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS CI F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS CI F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS CI F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS CI F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS CI F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS CI F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS CI F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS CI F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS CI F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS CI F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS CI F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS CI F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS CI F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS CI F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS CI F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS CI F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS CI F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS CI F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS CI F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS CI F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS CI F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS CI F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺

Comments: _____

LDC #: 16470A6
SDG #: See cover
VALIDATION FINDINGS WORKSHEET
Blanks

METHOD: Inorganics, Method See cover

Page: 1 of 1
Reviewer: MM
2nd Reviewer: R

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".
 N N/A Were all samples associated with a given method blank?
 N N/A Were any inorganic contaminants detected above the reporting limit in the method blanks? If yes, please see qualifications below.

Conc. units: ug/L Associated Samples: A1 (75X)

Analyte	Blank ID	Maximum ICB/CCB	Blank Action Limit	Sample Identification
SO4	MB	0.194	0.97	
Cl	0.220	0.189	1.1	

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
All contaminants within five times the method blank concentration were qualified as not detected, "N".

VALIDATION FINDINGS WORKSHEET
Initial and Continuing Calibration Calculation Verification

METHOD: Inorganics, Method See cover
 The correlation coefficient (r) for the calibration of TOC was recalculated. Calibration date: 12/31/06

An initial or continuing calibration verification percent recovery (%R) was recalculated for each type of analysis using the following formula:

$\%R = \frac{\text{Found}}{\text{True}} \times 100$ Where, Found = concentration of each analyte measured in the analysis of the ICV or CCV solution
 True = concentration of each analyte in the ICV or CCV source

Type of Analysis	Analyte	Conc. (mg/L) (units)	Found (units)	Recalculated		Reported	Acceptable (Y/N)
				r or %R	r or %R		
Initial calibration	Blank	0	0				
Calibration verification	Standard 1	1	944				
	Standard 2	10	6209				
	Standard 3	40	21870				Y
	Standard 4						
	Standard 5						
	Standard 6						
	Standard 7						
Calibration verification CCV	10	9.16		92	NR		Y
Calibration verification CCV	10	10.31		103	↓		↓
Calibration verification CCV	10	9.59		96	↓		↓

Comments: Refer to Calibration Verification findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

VALIDATION FINDINGS WORKSHEET
 Level IV Recalculation Worksheet

LDC #: 16490 AG
 SDG #: See cover

METHOD: Inorganics, Method See cover

Percent recoveries (%R) for a laboratory control sample and a matrix spike sample were recalculated using the following formula:

$$\%R = \frac{\text{Found}}{\text{True}} \times 100$$
 Where: Found = concentration of each analyte measured in the analysis of the sample. For the matrix spike calculation, Found = SSR (spiked sample result) - SR (sample result).
 True = concentration of each analyte in the source.

A sample and duplicate relative percent difference (RPD) was recalculated using the following formula:

$$RPD = \frac{|S-D|}{(S+D)/2} \times 100$$
 Where: S = Original sample concentration
 D = Duplicate sample concentration

Sample ID	Type of Analysis	Element	Found / S (units)	True / D (units)	Recalculated		Reported		Acceptable (Y/N)
					%R / RPD	%R / RPD			
<u>LES</u>	Laboratory control sample	<u>SO4</u>	<u>9.72</u>	<u>10</u>	<u>97</u>	<u>97</u>			<u>Y</u>
<u>TRK2314</u>	Matrix spike sample	<u>Br</u>	<u>5.67</u> (SSR-SR)	<u>5.0</u>	<u>113</u>	<u>113</u>			<u>Y</u>
<u>JPL0599</u>	Duplicate sample	<u>AlK</u>	<u>130</u>	<u>128</u>	<u>2</u>	<u>0</u>			<u>Y</u>

Comments: Refer to appropriate worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 16470AB
 SDG #: See cover

VALIDATION FINDINGS WORKSHEET
Sample Calculation Verification

Page: 1 of 1
 Reviewer: MM
 2nd reviewer: A

METHOD: Inorganics, Method See cover

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

- N N/A Have results been reported and calculated correctly?
- N N/A Are results within the calibrated range of the instruments?
- N N/A Are all detection limits below the CRQL?

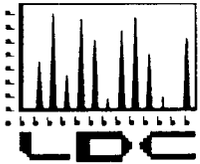
Compound (analyte) results for 1 reported with a positive detect were recalculated and verified using the following equation:

Concentration = _____ Recalculation:

$$NO_3 = \frac{\sqrt{4 \times 398.3587 \times (120357.12 + 634399) + 6186082} - 6186082}{2 \times 398.3587} = 17.495 \text{ mg/L}$$

#	Sample ID	Analyte	Reported Concentration (mg/L)	Calculated Concentration (mg/L)	Acceptable (Y/N)
1	1	Alk	200	200	Y
		Br	28	28	Y
		Cl	280	280	Y
		NO3	17	17	Y
		SO4	52	52	Y

Note: _____



LABORATORY DATA CONSULTANTS, INC.

7750 El Camino Real, Suite 2L Carlsbad, CA 92009 Phone: 760/634-0437 Fax: 760/634-0439

Tait Environmental Management, Inc.
701 N. Park Center Drive
Santa Ana, CA 92705
ATTN: Mr. Matt Hillman

May 1, 2007

SUBJECT: Boeing Realty Corp. Bldg C-1 Long Beach, Data Validation

Dear Mr. Hillman,

Enclosed are the final validation reports for the fraction listed below. These SDGs were received on April 9, 2007. Attachment 1 is a summary of the samples that were reviewed for each analysis.

LDC Project # 16591:

<u>SDG #</u>	<u>Fraction</u>
IQC0980, IQC1776	Volatiles, Semivolatiles, TPH as Extractables, Hexavalent Chromium

The data validation was performed under Tier 1, Tier 2 and Tier 3 guidelines. The analyses were validated using the following documents, as applicable to each method:

- USEPA, Contract Laboratory Program National Functional Guidelines for Organic Data Review, October 1999
- USEPA, Contract Laboratory Program National Functional Guidelines for Inorganic Data Review, October 1999
- EPA SW 846, Third Edition, Test Methods for Evaluating Solid Waste, update 1, July 1992; update IIA, August 1993; update II, September 1994; update IIB, January 1995; update III, December 1996; update IIIA, April 1998

Please feel free to contact us if you have any questions.

Sincerely,

Stella S. Cuenco
Project Manager/Senior Chemist

**Boeing Realty Corp., Bldg C-1 Long Beach
Data Validation Reports
LDC# 16591**

Volatiles

LDC

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Boeing Realty Corp., Bldg C-1 Long Beach

Collection Date: March 15, 2007

LDC Report Date: April 30, 2007

Matrix: Water

Parameters: Volatiles

Validation Level: Tier 3

Laboratory: TestAmerica

Sample Delivery Group (SDG): IQC1776

Sample Identification

MW3017_WG031507_0001

Introduction

This data review covers one water sample listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 8260B for Volatiles.

This review follows a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Organic Data Review (October 1999) as there are no current guidelines for the method stated above.

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

Blank results are summarized in Section V.

Field duplicates are summarized in Section XVI.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- N Presumptive evidence of presence of the constituent.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. GC/MS Instrument Performance Check

Instrument performance was checked at 12 hour intervals.

All ion abundance requirements were met.

III. Initial Calibration

Initial calibration was performed using required standard concentrations.

Percent relative standard deviations (%RSD) were less than or equal to 15.0% for each individual compound and less than or equal to 30.0% for calibration check compounds (CCCs).

In the case where %RSD was greater than 15.0%, the laboratory used a calibration curve to evaluate the compound. All coefficients of determination (r^2) were greater than or equal to 0.990 .

For the purposes of technical evaluation, all compounds were evaluated against the 30.0% (%RSD) National Functional Guideline criteria. Unless noted above, all compounds were within the validation criteria.

Average relative response factors (RRF) for all volatile target compounds and system performance check compounds (SPCCs) were within method and validation criteria with the following exceptions:

Date	Compound	RRF (Limits)	Associated Samples	Flag	A or P
2/28/07	2-Butanone	0.037 (≥ 0.05)	All samples in SDG IQC1776	J (all detects) UJ (all non-detects)	A

IV. Continuing Calibration

Continuing calibration was performed at the required frequencies.

Percent differences (%D) between the initial calibration RRF and the continuing calibration RRF were within the method criteria of less than or equal to 20.0% for calibration check compounds (CCCs).

For the purposes of technical evaluation, all compounds were evaluated against the 25.0% (%D) National Functional Guideline criteria. Unless noted above, all compounds were within the validation criteria.

All of the continuing calibration RRF values were within method and validation criteria with the following exceptions:

Date	Compound	RRF (Limits)	Associated Samples	Flag	A or P
3/20/07	2-Butanone	0.040 (≥ 0.05)	All samples in SDG IQC1776	J (all detects) UJ (all non-detects)	A

V. Blanks

Method blanks were reviewed for each matrix as applicable. No volatile contaminants were found in the method blanks with the following exceptions:

Method Blank ID	Analysis Date	Compound TIC (RT in minutes)	Concentration	Associated Samples
7C20023-BLK1	3/20/07	Tetrahydrofuran	8.39 ug/L	All samples in SDG IQC1776

Sample concentrations were compared to concentrations detected in the method blanks. The sample concentrations were either not detected or were significantly greater ($>10X$ for common contaminants, $>5X$ for other contaminants) than the concentrations found in the associated method blanks.

VI. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VII. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were not within QC limits. Since there were no associated samples, no data were qualified.

VIII. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) were within QC limits with the following exceptions:

LCS ID	Compound	%R (Limits)	Associated Samples	Flag	A or P
7C20023-BS1	2-Butanone 2-Hexanone	160 (40-140) 151 (45-140)	All samples in SDG IQC1776	J (all detects) J (all detects)	P

IX. Regional Quality Assurance and Quality Control

Not applicable.

X. Internal Standards

All internal standard areas and retention times were within QC limits.

XI. Target Compound Identifications

All target compound identifications were within validation criteria.

XII. Compound Quantitation and CRQLs

All compound quantitation and CRQLs were within validation criteria.

XIII. Tentatively Identified Compounds (TICs)

Tentatively identified compounds were not reported by the laboratory.

XIV. System Performance

The system performance was acceptable.

XV. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

XVI. Field Duplicates

No field duplicates were identified in this SDG.

XVII. Field Blanks

No field blanks were identified in this SDG.

**Boeing Realty Corp., Bldg C-1 Long Beach
 Volatiles - Data Qualification Summary - SDG IQC1776**

SDG	Sample	Compound	Flag	A or P	Reason
IQC1776	MW3017_WG031507_0001	2-Butanone	J (all detects) UJ (all non-detects)	A	Initial calibration (RRF)
IQC1776	MW3017_WG031507_0001	2-Butanone	J (all detects) UJ (all non-detects)	A	Continuing calibration (RRF)
IQC1776	MW3017_WG031507_0001	2-Butanone 2-Hexanone	J (all detects) J (all detects)	P	Laboratory control samples (%R)

**Boeing Realty Corp., Bldg C-1 Long Beach
 Volatiles - Laboratory Blank Data Qualification Summary - SDG IQC1776**

No Sample Data Qualified in this SDG

TestAmerica

ANALYTICAL TESTING CORPORATION

17461 Derian Avenue, Suite 100, Irvine, CA 92614 (949) 261-1022 Fax: (949) 260-3297

TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-1 Long Beach
EM-2701
Report Number: IQC1776

Sampled: 03/15/07
Received: 03/15/07

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC1776-04 (MW3017_WG031507_0001 - Water)									
Reporting Units: ug/l									
Benzene	EPA 8260B	7C20023	0.28	1.0	1.1	1	03/20/07	03/20/07	
Bromobenzene	EPA 8260B	7C20023	0.27	1.0	ND	1	03/20/07	03/20/07	
Bromochloromethane	EPA 8260B	7C20023	0.32	1.0	ND	1	03/20/07	03/20/07	
Bromodichloromethane	EPA 8260B	7C20023	0.30	1.0	ND	1	03/20/07	03/20/07	
Bromoform	EPA 8260B	7C20023	0.40	1.0	ND	1	03/20/07	03/20/07	
Bromomethane	EPA 8260B	7C20023	0.42	1.0	ND	1	03/20/07	03/20/07	
2-Butanone (MEK)	EPA 8260B	7C20023	3.8	5.0	ND	1	03/20/07	03/20/07	L
n-Butylbenzene	EPA 8260B	7C20023	0.37	1.0	ND	1	03/20/07	03/20/07	
sec-Butylbenzene	EPA 8260B	7C20023	0.25	1.0	8.3	1	03/20/07	03/20/07	
tert-Butylbenzene	EPA 8260B	7C20023	0.22	1.0	0.89	1	03/20/07	03/20/07	J
Carbon Disulfide	EPA 8260B	7C20023	0.48	1.0	0.68	1	03/20/07	03/20/07	J
Carbon tetrachloride	EPA 8260B	7C20023	0.28	0.50	ND	1	03/20/07	03/20/07	
Chlorobenzene	EPA 8260B	7C20023	0.36	1.0	ND	1	03/20/07	03/20/07	
Chloroethane	EPA 8260B	7C20023	0.40	2.0	0.65	1	03/20/07	03/20/07	J
Chloroform	EPA 8260B	7C20023	0.33	1.0	ND	1	03/20/07	03/20/07	
Chloromethane	EPA 8260B	7C20023	0.40	2.0	ND	1	03/20/07	03/20/07	
2-Chlorotoluene	EPA 8260B	7C20023	0.28	1.0	ND	1	03/20/07	03/20/07	
4-Chlorotoluene	EPA 8260B	7C20023	0.29	1.0	ND	1	03/20/07	03/20/07	
1,2-Dibromo-3-chloropropane	EPA 8260B	7C20023	0.97	2.0	ND	1	03/20/07	03/20/07	
Dibromochloromethane	EPA 8260B	7C20023	0.28	1.0	ND	1	03/20/07	03/20/07	
1,2-Dibromoethane (EDB)	EPA 8260B	7C20023	0.40	1.0	ND	1	03/20/07	03/20/07	
1,4-Dichlorobenzene	EPA 8260B	7C20023	0.37	1.0	ND	1	03/20/07	03/20/07	
1,2-Dichlorobenzene	EPA 8260B	7C20023	0.32	1.0	ND	1	03/20/07	03/20/07	
1,3-Dichlorobenzene	EPA 8260B	7C20023	0.35	1.0	ND	1	03/20/07	03/20/07	
Dichlorodifluoromethane	EPA 8260B	7C20023	0.79	1.0	ND	1	03/20/07	03/20/07	
1,2-Dichloroethane	EPA 8260B	7C20023	0.28	0.50	1.2	1	03/20/07	03/20/07	
1,1-Dichloroethane	EPA 8260B	7C20023	0.27	1.0	2.6	1	03/20/07	03/20/07	
1,1-Dichloroethene	EPA 8260B	7C20023	0.42	1.0	ND	1	03/20/07	03/20/07	
cis-1,2-Dichloroethene	EPA 8260B	7C20023	0.32	1.0	ND	1	03/20/07	03/20/07	
trans-1,2-Dichloroethene	EPA 8260B	7C20023	0.27	1.0	ND	1	03/20/07	03/20/07	
1,2-Dichloropropane	EPA 8260B	7C20023	0.35	1.0	ND	1	03/20/07	03/20/07	
2,2-Dichloropropane	EPA 8260B	7C20023	0.34	1.0	ND	1	03/20/07	03/20/07	
cis-1,3-Dichloropropene	EPA 8260B	7C20023	0.22	0.50	ND	1	03/20/07	03/20/07	
1,1-Dichloropropene	EPA 8260B	7C20023	0.28	1.0	ND	1	03/20/07	03/20/07	
trans-1,3-Dichloropropene	EPA 8260B	7C20023	0.32	0.50	ND	1	03/20/07	03/20/07	
Ethylbenzene	EPA 8260B	7C20023	0.25	1.0	9.4	1	03/20/07	03/20/07	
Hexachlorobutadiene	EPA 8260B	7C20023	0.38	1.0	ND	1	03/20/07	03/20/07	
2-Hexanone	EPA 8260B	7C20023	2.6	6.0	ND	1	03/20/07	03/20/07	L
Iodomethane	EPA 8260B	7C20023	1.0	2.0	ND	1	03/20/07	03/20/07	
Isopropylbenzene	EPA 8260B	7C20023	0.25	1.0	8.8	1	03/20/07	03/20/07	
p-Isopropyltoluene	EPA 8260B	7C20023	0.28	1.0	1.6	1	03/20/07	03/20/07	

TestAmerica - Irvine, CA
Nicholas Marz
Project Manager

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IQC1776 <Page 9 of 37>

TestAmerica

ANALYTICAL TESTING CORPORATION

17461 Derian Avenue, Suite 100, Irvine, CA 92614 (949) 261-1022 Fax: (949) 260-3297

TAIT Environmental/Boeing 701 N. Parkcenter Drive Santa Ana, CA 92705 Attention: Mehmet Pehlivan	Project ID: Boeing C-1 Long Beach EM-2701 Report Number: IQC1776	Sampled: 03/15/07 Received: 03/15/07
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VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC1776-04 (MW3017_WG031507_0001 - Water) - cont.									
Reporting Units: ug/l									
Methyl-tert-butyl Ether (MTBE)	EPA 8260B	7C20023	0.32	1.0	ND	1	03/20/07	03/20/07	
Methylene chloride	EPA 8260B	7C20023	0.95	1.0	ND	1	03/20/07	03/20/07	
4-Methyl-2-pentanone (MIBK)	EPA 8260B	7C20023	3.5	5.0	ND	1	03/20/07	03/20/07	
n-Propylbenzene	EPA 8260B	7C20023	0.27	1.0	9.4	1	03/20/07	03/20/07	
Styrene	EPA 8260B	7C20023	0.16	1.0	ND	1	03/20/07	03/20/07	
1,1,1,2-Tetrachloroethane	EPA 8260B	7C20023	0.27	1.0	ND	1	03/20/07	03/20/07	
1,1,2,2-Tetrachloroethane	EPA 8260B	7C20023	0.24	1.0	ND	1	03/20/07	03/20/07	
Tetrachloroethene	EPA 8260B	7C20023	0.32	1.0	ND	1	03/20/07	03/20/07	
Tetrahydrofuran (THF)	EPA 8260B	7C20023	3.5	10	ND	1	03/20/07	03/20/07	
Toluene	EPA 8260B	7C20023	0.36	1.0	ND	1	03/20/07	03/20/07	
1,2,3-Trichlorobenzene	EPA 8260B	7C20023	0.30	1.0	ND	1	03/20/07	03/20/07	
1,2,4-Trichlorobenzene	EPA 8260B	7C20023	0.48	1.0	ND	1	03/20/07	03/20/07	
1,1,2-Trichloroethane	EPA 8260B	7C20023	0.30	1.0	ND	1	03/20/07	03/20/07	
1,1,1-Trichloroethane	EPA 8260B	7C20023	0.30	1.0	ND	1	03/20/07	03/20/07	
Trichloroethene	EPA 8260B	7C20023	0.26	1.0	ND	1	03/20/07	03/20/07	
Trichlorofluoromethane	EPA 8260B	7C20023	0.34	2.0	ND	1	03/20/07	03/20/07	
1,2,3-Trichloropropane	EPA 8260B	7C20023	0.40	1.0	ND	1	03/20/07	03/20/07	
1,2,4-Trimethylbenzene	EPA 8260B	7C20023	0.23	1.0	8.3	1	03/20/07	03/20/07	
1,3,5-Trimethylbenzene	EPA 8260B	7C20023	0.26	1.0	2.5	1	03/20/07	03/20/07	
Vinyl acetate	EPA 8260B	7C20023	1.7	6.0	ND	1	03/20/07	03/20/07	
Vinyl chloride	EPA 8260B	7C20023	0.30	0.50	ND	1	03/20/07	03/20/07	
Xylenes, Total	EPA 8260B	7C20023	0.90	1.0	1.9	1	03/20/07	03/20/07	
Surrogate: 4-Bromofluorobenzene (80-120%)					111 %				
Surrogate: Dibromofluoromethane (80-120%)					107 %				
Surrogate: Toluene-d8 (80-120%)					107 %				

TestAmerica - Irvine, CA
Nicholas Marz
Project Manager

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ANALYTICAL TESTING CORPORATION

17461 Deegan Avenue, Suite 100, Irvine, CA 92614 (949) 261-1022 Fax: (949) 260-3297

TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-1 Long Beach
EM-2701
Report Number: IQC1776

Sampled: 03/15/07
Received: 03/15/07

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC1776-04RE1 (MW3017_WG031507_0001 - Water) - cont.									
Reporting Units: ug/l									
Acetone	EPA 8260B	7C22010	4.5	10	ND	1	03/22/07	03/22/07	
Surrogate: 4-Bromofluorobenzene (80-120%)					100 %				
Surrogate: Dibromofluoromethane (80-120%)					96 %				
Surrogate: Toluene-d8 (80-120%)					98 %				

TestAmerica - Irvine, CA
Nicholas Marz
Project Manager

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IQC1776 <Page 11 of 37>

LDC #: 16591B1
 SDG #: IQC1776
 Laboratory: Test America

VALIDATION COMPLETENESS WORKSHEET
 Tier 3

Date: 4/25/07
 Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: GC/MS Volatiles (EPA SW 846 Method 8260B)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: 3/15/07
II.	GC/MS Instrument performance check	A	
III.	Initial calibration	SW	% RSD, r ² 20.990
IV.	Continuing calibration	SW	
V.	Blanks	SW	
VI.	Surrogate spikes	A	
VII.	Matrix spike/Matrix spike duplicates	SW	MW301B_WG031507_0001
VIII.	Laboratory control samples	SW	LCs
IX.	Regional Quality Assurance and Quality Control	N	
X.	Internal standards	A	
XI.	Target compound identification	A	
XII.	Compound quantitation/CRQLs	A	
XIII.	Tentatively identified compounds (TICs)	N	not reported
XIV.	System performance	A	
XV.	Overall assessment of data	A	
XVI.	Field duplicates	N	
XVII.	Field blanks	N	

Note: A = Acceptable ND = No compounds detected D = Duplicate
 N = Not provided/applicable R = Rinsate TB = Trip blank
 SW = See worksheet FB = Field blank EB = Equipment blank

Validated Samples: wait

1	MW3017-WG031507_0001	11	7C20023-BLK1	21		31	
2	MW3017-WG031507_0001MSD	12	7C22010-BLK1	22		32	
3	MW3017-WG031507_0001MSD	13		23		33	
4		14		24		34	
5		15		25		35	
6		16		26		36	
7		17		27		37	
8		18		28		38	
9		19		29		39	
10		20		30		40	

DC #: 16591 B1
 SDG #: see cover

VALIDATION FINDINGS CHECKLIST

Page: 1 of 2
 Reviewer: FR
 2nd Reviewer: FR

Method: Volatiles (EPA SW 846 Method 8260B)

Validation Area	Yes	No	NA	Findings/Comments
All technical holding times were met.	/			
Cooler temperature criteria was met.	/			
Were the BFB performance results reviewed and found to be within the specified criteria?	/			
Were all samples analyzed within the 12 hour clock criteria?	/			
Did the laboratory perform a 5 point calibration prior to sample analysis?	/			
Were all percent relative standard deviations (%RSD) and relative response factors (RRF) within method criteria for all CCCs and SPCCs?	/			
Was a curve fit used for evaluation?	/			
Did the initial calibration meet the curve fit acceptance criteria of > 0.990?	/			
Were all percent relative standard deviations (%RSD) ≤ 30% and relative response factors (RRF) > 0.05?		/		
Was a continuing calibration standard analyzed at least once every 12 hours for each instrument?	/			
Were all percent differences (%D) and relative response factors (RRF) within method criteria for all CCCs and SPCCs?	/			
Were all percent differences (%D) ≤ 25% and relative response factors (RRF) ≥ 0.05?		/		
Was a method blank associated with every sample in this SDG?	/			
Was a method blank analyzed at least once every 12 hours for each matrix and concentration?	/			
Was there contamination in the method blanks? If yes, please see the Blanks validation completeness worksheet.		/		
Were all surrogate %R within QC limits?	/			
If the percent recovery (%R) for one or more surrogates was out of QC limits, was a reanalysis performed to confirm samples with %R outside of criteria?			/	
Were a matrix spike (MS) and matrix spike duplicate (MSD) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD. Soil / Water.	/			
Was a MS/MSD analyzed every 20 samples of each matrix?	/			
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the QC limits?		/		
Was an LCS analyzed for this SDG?	/			

DC #: 16591B1
 S #: see cover

VALIDATION FINDINGS CHECKLIST

Page: 2 of 2
 Reviewer: JS
 2nd Reviewer: RL

Validation Area	Yes	No	NA	Findings/Comments
Was an LCS analyzed per analytical batch?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the QC limits?	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
Were performance evaluation (PE) samples performed?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were the performance evaluation (PE) samples within the acceptance limits?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were internal standard area counts within -50% or +100% of the associated calibration standard?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were retention times within + 30 seconds of the associated calibration standard?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were relative retention times (RRT's) within + 0.06 RRT units of the standard?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Did compound spectra meet specified EPA "Functional Guidelines" criteria?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were chromatogram peaks verified and accounted for?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the correct internal standard (IS), quantitation ion and relative response factor (RRF) used to quantitate the compound?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were compound quantitation and CRQLs adjusted to reflect all sample dilutions and dry weight factors applicable to level IV validation?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the major ions (> 10 percent relative intensity) in the reference spectrum evaluated in sample spectrum?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were relative intensities of the major ions within ± 20% between the sample and the reference spectra?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Did the raw data indicate that the laboratory performed a library search for all required peaks in the chromatograms (samples and blanks)?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
System performance was found to be acceptable.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Overall assessment of data was found to be acceptable.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Field duplicate pairs were identified in this SDG.	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
Target compounds were detected in the field duplicates.	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Field blanks were identified in this SDG.	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
Target compounds were detected in the field blanks.	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	

TARGET COMPOUND WORKSHEET

METHOD: VOA (EPA SW 846 Method 8260B)

A. Chloromethane*	S. Trichloroethene	KK. Trichlorofluoromethane	CCC. tert-Butylbenzene	UUU. 1,2-Dichlorotetrafluoroethane
B. Bromomethane	T. Dibromochloromethane	LL. Methyl-tert-butyl ether	DDD. 1,2,4-Trimethylbenzene	VVV. 4-Ethyltoluene
C. Vinyl chloride**	U. 1,1,2-Trichloroethane	MM. 1,2-Dibromo-3-chloropropane	EEE. sec-Butylbenzene	WWW. Ethanol
D. Chloroethane	V. Benzene	NN. Methyl ethyl ketone	FFF. 1,3-Dichlorobenzene	XXX. Diisopropyl ether
E. Methylene chloride	W. trans-1,3-Dichloropropane	OO. 2,2-Dichloropropane	GGG. p-Isopropyltoluene	YYY. tert-Butanol
F. Acetone	X. Bromoform*	PP. Bromochloromethane	HHH. 1,4-Dichlorobenzene	ZZZ. tert-Butyl alcohol
G. Carbon disulfide	Y. 4-Methyl-2-pentanone	QQ. 1,1-Dichloropropane	III. n-Butylbenzene	AAA. Ethyl tert-butyl ether
H. 1,1-Dichloroethane**	Z. 2-Hexanone	RR. Dibromomethane	JJJ. 1,2-Dichlorobenzene	BBB. tert-Amyl methyl ether
I. 1,1-Dichloroethane*	AA. Tetrachloroethene	SS. 1,3-Dichloropropane	KKK. 1,2,4-Trichlorobenzene	CCCC. 1-Chlorohexane
J. 1,2-Dichloroethane, total	BB. 1,1,2,2-Tetrachloroethane*	TT. 1,2-Dibromoethane	LLL. Hexachlorobutadiene	DDDD. Isopropyl alcohol
K. Chloroform**	CC. Toluene**	UU. 1,1,1,2-Tetrachloroethane	MMM. Naphthalene	EEEE. Acetonitrile
L. 1,2-Dichloroethane	DD. Chlorobenzene*	VV. Isopropylbenzene	NNN. 1,2,3-Trichlorobenzene	FFFF. Acrolein
M. 2-Butanone	EE. Ethylbenzene**	WW. Bromobenzene	OOO. 1,3,5-Trichlorobenzene	GGGG. Acrylonitrile
N. 1,1,1-Trichloroethane	FF. Styrene	XX. 1,2,3-Trichloropropane	PPP. trans-1,2-Dichloroethane	HHHH. 1,4-Dioxane
O. Carbon tetrachloride	GG. Xylenes, total	YY. n-Propylbenzene	QQQ. cis-1,2-Dichloroethane	IIII. Isobutyl alcohol
P. Bromodichloromethane	HH. Vinyl acetate	ZZ. 2-Chlorotoluene	RRR. m,p-Xylenes	JJJJ. Methacrylonitrile
Q. 1,2-Dichloropropane**	II. 2-Chloroethylvinyl ether	AAA. 1,3,5-Trimethylbenzene	SSS. o-Xylene	KKKK. Propionitrile
R. cis-1,3-Dichloropropane	JJ. Dichlorodifluoromethane	BBB. 4-Chlorotoluene	TTT. 1,1,2-Trichloro-1,2,2-trifluoroethane	LLLL

* = System performance check compounds (SPCC) for RRF ; ** = Calibration check compounds (CCC) for %RSD.

LDC #: 10511B)

SDG #: see cover

VALIDATION FINDINGS WORKSHEET

Initial Calibration

Page: 1 of 1
Reviewer: B
2nd Reviewer: A

METHOD: GC/MS VOA (EPA SW 846 Method 8260B)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

- Y N N/A Did the laboratory perform a 5 point calibration prior to sample analysis?
- Y N N/A Were percent relative standard deviations (%RSD) and relative response factors (RRF) within method criteria for all CCC's and SPCC's?
- Y N N/A Was a curve fit used for evaluation? If yes, what was the acceptance criteria used for evaluation? _____
- Y N N/A Did the initial calibration meet the acceptance criteria?
- Y N N/A Were all %RSDs and RRFs within the validation criteria of ≤ 30 %RSD and ≥ 0.05 RRF ?

#	Date	Standard ID	Compound	Finding %RSD (Limit: $\leq 30.0\%$)	Finding RRF (Limit: ≥ 0.05)	Associated Samples	Qualifications
	2/24/07	ICAL	M		0.037	All 1014 + 7C20023-B14	JUJ/A

LDC #: 1691 B1
SDG #: F 601776

VALIDATION FINDINGS WORKSHEET
Blanks

Page: 1 of 1
Reviewer: AK
2nd Reviewer: AK

METHOD: GC/MS VOA (EPA SW 846 Method 8260B)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

N N/A Was a method blank associated with every sample in this SDG?

Y N/A Was a method blank analyzed at least once every 12 hours for each matrix and concentration?

Y N/A Was there contamination in the method blanks? If yes, please see the qualifications below.

Blank analysis date: 3/20/07

Conc. units: ug/l

Associated Samples: # 1 (ND)

Compound	Blank ID	Sample Identification
Tetrahydrofuran	TC20023- 8.39	BUK 1
Methylene chloride		
Acetone		
CRQL		

Blank analysis date: _____

Conc. units: _____

Associated Samples: _____

Compound	Blank ID	Sample Identification
Tetrahydrofuran		
Methylene chloride		
Acetone		
CRQL		

All results were qualified using the criteria stated below except those circled.

Note: Common contaminants such as Methylene chloride, Acetone, 2-Butanone, Carbon disulfide and TICs that were detected in samples within ten times the associated method blank concentration were qualified as not detected, "U". Other contaminants within five times the method blank concentration were also qualified as not detected, "U".

LDC #: 16591B
 SDG #: 160176

VALIDATION FINDINGS WORKSHEET
Initial Calibration Calculation Verification

Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: GC/MS VOA (EPA SW 846 Method 8260B)

The Relative Response Factor (RRF), average RRF, and percent relative standard deviation (%RSD) were recalculated for the compounds identified below using the following calculations:

$RRF = (A_i/C_i)/(A_s/C_s)$
 average RRF = sum of the RRFs/number of standards
 $\%RSD = 100 * (S/X)$
 A_i = Area of compound,
 C_i = Concentration of compound,
 S = Standard deviation of the RRFs
 X = Mean of the RRFs
 A_s = Area of associated internal standard
 C_s = Concentration of internal standard

#	Standard ID	Calibration Date	Compound (Reference Internal Standard)	Reported		Recalculated		Reported		Recalculated	
				RRF (10 std)	RRF (10 std)	Average RRF (Initial)	Average RRF (Initial)	%RSD	%RSD		
1	1CAL	2/28/07	Methylene chloride (1st internal standard)	0.436	0.436	0.474	0.474	7.78	7.78		
			Trichlorethene (2nd internal standard)	0.314	0.314	0.321	0.331	5.92	5.92		
			Toluene (3rd internal standard)	0.269	0.269	0.281	0.281	5.18	5.18		
			1,2-DCB	1.306	1.306	1.314	1.314	10.24	10.24		
2	1CAL	2/28/07	Methylene chloride (1st internal standard)	0.169	0.169	0.179	0.179	22.59	22.59		
			Trichlorethene (2nd internal standard)								
			Toluene (3rd internal standard)								
			Methylene chloride (1st internal standard)								
3	1CAL	2/28/07	Methylene chloride (1st internal standard)								
			Trichlorethene (2nd internal standard)								
			Toluene (3rd internal standard)								
			Methylene chloride (1st internal standard)								
4	1CAL	2/28/07	Methylene chloride (1st internal standard)								
			Trichlorethene (2nd internal standard)								
			Toluene (3rd internal standard)								
			Methylene chloride (1st internal standard)								

Comments: Refer to Initial Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 16591B
 SDG #: for cover

VALIDATION FINDINGS WORKSHEET
Continuing Calibration Results Verification

Page: 1 of 1
 Reviewer: RR
 2nd Reviewer: RR

METHOD: GC/MS VOA (EPA SW 846 Method 8260B)

The percent difference (%D) of the initial calibration average Relative Response Factors (RRFs) and the continuing calibration RRFs were recalculated for the compounds identified below using the following calculation:

% Difference = $100 \cdot (\text{ave. RRF} - \text{RRF}) / \text{ave. RRF}$
 RRF = $(A_x)(C_s) / (A_s)(C_x)$

Where: ave. RRF = initial calibration average RRF
 RRF = continuing calibration RRF
 A_x = Area of compound,
 C_x = Concentration of compound,
 A_s = Area of associated internal standard
 C_s = Concentration of internal standard

#	Standard ID	Calibration Date	Compound (Reference Internal Standard)	Average RRF (Initial)	Reported		Recalculated	
					RRF (CC)	%D	RRF (CC)	%D
1	ceV	3/20/07	Methylene chloride (1st internal standard)	0.474	0.492	3.8	0.492	3.8
	6.19PM		Trichlorethene (2nd internal standard)	0.331	0.339	2.4	0.339	2.4
			Toluene (3rd internal standard)	0.281	0.281	0	0.281	0
2			1,2-DCB	1.314	1.334	1.5	1.334	1.5
			Methylene chloride (1st internal standard)					
			Trichlorethene (2nd internal standard)					
3	ceV	3/22/07	Toluene (3rd internal standard)	0.179	0.207	15.6	0.207	15.6
	7:10 AM		Methylene chloride (1st internal standard)					
			Trichlorethene (2nd internal standard)					
4			Toluene (3rd internal standard)					
			Methylene chloride (1st internal standard)					
			Trichlorethene (2nd internal standard)					

Comments: Refer to Continuing Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

CONCL:15B

LDC #: 16591B
 SDG #: see cover

VALIDATION FINDINGS WORKSHEET
Surrogate Results Verification

Page: 1 of 1
 Reviewer: [Signature]
 2nd reviewer: [Signature]

METHOD: GC/MS VOA (EPA SW 846 Method 8260B)

The percent recoveries (%R) of surrogates were recalculated for the compounds identified below using the following calculation:

% Recovery: $SF/SS * 100$

Where: SF = Surrogate Found
 SS = Surrogate Spiked

Sample ID: #1

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Toluene-d8	25.0	26.85	107	107	0
Bromofluorobenzene	↓	27.67	111	111	↓
1,2-Dichloroethane-d4					
Dibromofluoromethane	↓	26.84	107	107	↓

Sample ID: _____

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Toluene-d8					
Bromofluorobenzene					
1,2-Dichloroethane-d4					
Dibromofluoromethane					

Sample ID: _____

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Toluene-d8					
Bromofluorobenzene					
1,2-Dichloroethane-d4					
Dibromofluoromethane					

Sample ID: _____

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Toluene-d8					
Bromofluorobenzene					
1,2-Dichloroethane-d4					
Dibromofluoromethane					

Sample ID: _____

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Toluene-d8					
Bromofluorobenzene					
1,2-Dichloroethane-d4					
Dibromofluoromethane					

LDC #: 16591B
 SDG #: per comp

VALIDATION FINDINGS WORKSHEET
Matrix Spike/Matrix Spike Duplicates Results Verification

Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: GC/MS VOA (EPA SW 846 Method 8260B)

The percent recoveries (%R) and Relative Percent Difference (RPD) of the matrix spike and matrix spike duplicate were recalculated for the compounds identified below using the following calculation:

$\% \text{ Recovery} = 100 * (\text{SSC} - \text{SC}) / \text{SA}$ Where: SSC = Spiked sample concentration SC = Sample concentration
 SA = Spike added

$\text{RPD} = 1 \text{ MSC} - \text{MSDC} / 2(\text{MSC} + \text{MSDC})$ MSC = Matrix spike percent recovery MSDC = Matrix spike duplicate percent recovery

MS/MSD sample: MW3018 - W4031507 - 0001

Compound	Spike Added (ug/L)		Sample Concentration (ug/L)	Spiked Sample Concentration (ug/L)		Matrix Spike Percent Recovery		Matrix Spike Duplicate Percent Recovery		MS/MSD RPD	
	MS	MSD		MS	MSD	Reported	Recalc.	Reported	Recalc.	Reported	Recalculated
1,1-Dichloroethene	25.0	25.0	ND	24.7	24.4	99	99	98	98	1	1
Trichloroethene	↓		ND	25.7	25.4	103	103	102	102	0	0
Benzene	25.0		0.94	27.9	27.7	108	108	107	107	1	1
Toluene	↓		ND	27.0	26.9	108	108	108	108	0	0
Chlorobenzene	↓		ND	27.6	27.3	110	110	109	109	1	1

Comments: Refer to Matrix Spike/Matrix Spike Duplicates findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 16591B
 SDG #: file cover

VALIDATION FINDINGS WORKSHEET
Laboratory Control Sample Results Verification

Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: GC/MS VOA (EPA SW 846 Method 8260B)

The percent recoveries (%R) and Relative Percent Difference (RPD) of the laboratory control sample and laboratory control sample duplicate (if applicable) were recalculated for the compounds identified below using the following calculation:

% Recovery = $100 * SSC/SA$

Where: SSC = Spiked sample concentration
 SA = Spike added

RPD = $1 LCS - LCSD \mid * 2 / (LCS + LCSD)$

LCS = Laboratory control sample percent recovery

LCSD = Laboratory control sample duplicate percent recovery

LCS ID: 7220023 - B2

Compound	Spike Added (ug/L)		Spiked Sample Concentration (ug/L)		LCS Percent Recovery		LCSD Percent Recovery		LCS/LCSD RPD	
	LCS	LCSD	LCS	LCSD	Reported	Recalc.	Reported	Recalc.	Reported	Recalculated
1,1-Dichloroethene	25.0	NA	23.8	NA	95	95	106	106		
Trichloroethene			26.4		110	106				
Benzene			27.4		109	110				
Toluene			27.2		111	109				
Chlorobenzene			27.7			111			NA	

Comments: Refer to Laboratory Control Sample findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

DC #: 16591B1
 IG #: see cover

VALIDATION FINDINGS WORKSHEET

Sample Calculation Verification

Page: 1 of 1
 Reviewer: R
 2nd reviewer: R

METHOD: GC/MS VOA (EPA SW 846 Method 8260B)

- Y/N N/A Were all reported results recalculated and verified for all level IV samples?
- Y/N N/A Were all recalculated results for detected target compounds agree within 10.0% of the reported results?

$$\text{Concentration} = \frac{(A_s)(I)(DF)}{(A_c)(RRF)(V_s)(\%S)}$$

- A_s = Area of the characteristic ion (EICP) for the compound to be measured
- A_c = Area of the characteristic ion (EICP) for the specific internal standard
- I = Amount of internal standard added in nanograms (ng)
- RRF = Relative response factor of the calibration standard.
- V_s = Volume or weight of sample pruged in milliliters (ml) or grams (g).
- Df = Dilution factor.
- %S = Percent solids, applicable to soils and solid matrices only.

Example:

Sample I.D. # 1 Benzene

$$\text{Conc.} = \frac{(69826)(25)}{(14403)(1.069)} = 1.13 \text{ ug/L}$$

#	Sample ID	Compound	Reported Concentration ()	Calculated Concentration ()	Qualification

**Boeing Realty Corp., Bldg C-1 Long Beach
Data Validation Reports
LDC# 16591**

Semivolatiles

LDC

LDC Report# 16591B2

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Boeing Realty Corp., Bldg C-1 Long Beach
Collection Date: March 15, 2007
LDC Report Date: April 30, 2007
Matrix: Water
Parameters: Semivolatiles
Validation Level: Tier 3
Laboratory: TestAmerica

Sample Delivery Group (SDG): IQC1776

Sample Identification

MW3017_WG031507_0001

Introduction

This data review covers one water sample listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 8270C for Semivolatiles.

This review follows a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Organic Data Review (October 1999) as there are no current guidelines for the method stated above.

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified a P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

Blank results are summarized in Section V.

Field duplicates are summarized in Section XVI.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- N Presumptive evidence of presence of the constituent.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. GC/MS Instrument Performance Check

Instrument performance was checked at 12 hour intervals.

All ion abundance requirements were met.

III. Initial Calibration

Initial calibration was performed using required standard concentrations.

Percent relative standard deviations (%RSD) were less than or equal to 15.0% for each individual compound and less than or equal to 30.0% for calibration check compounds (CCCs).

In the case where %RSD was greater than 15.0%, the laboratory used a calibration curve to evaluate the compound. All coefficients of determination (r^2) were greater than or equal to 0.990 .

For the purposes of technical evaluation, all compounds were evaluated against the 30.0% (%RSD) National Functional Guideline criteria. Unless noted above, all compounds were within the validation criteria.

Average relative response factors (RRF) for all semivolatile target compounds and system performance check compounds (SPCCs) were greater than or equal to 0.05 as required.

IV. Continuing Calibration

Continuing calibration was performed at the required frequencies.

Percent differences (%D) between the initial calibration RRF and the continuing calibration RRF were within the method criteria of less than or equal to 20.0% for calibration check compounds (CCCs).

For the purposes of technical evaluation, all compounds were evaluated against the 25.0% (%D) National Functional Guideline criteria. Unless noted above, all compounds were within the validation criteria.

The percent difference (%D) of the second source calibration standard were less than or equal to 25.0% for all compounds.

All of the continuing calibration RRF values were greater than or equal to 0.05 .

V. Blanks

Method blanks were reviewed for each matrix as applicable. No semivolatile contaminants were found in the method blanks.

VI. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VII. Matrix Spike/Matrix Spike Duplicates

The laboratory has indicated that there were no matrix spike (MS) and matrix spike duplicate (MSD) analyses specified for the samples in this SDG, and therefore matrix spike and matrix spike duplicate analyses were not performed for this SDG.

VIII. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the following exceptions:

LCS ID (Associated Samples)	Compound	LCS %R (Limits)	LCSD %R (Limits)	RPD (Limits)	Flag	A or P
7C16066-LCS/D (All samples in SDG IQC1776)	1,3-Dichlorobenzene	29 (35-120)	-	34 (≤25)	J (all detects) UJ (all non-detects)	P
	1,4-Dichlorobenzene	34 (35-120)	-	27 (≤25)		
	1,2-Dichlorobenzene	35 (40-120)	-	31 (≤25)		
	Hexachlorobutadiene	37 (40-120)	-	32 (≤25)		
	Hexachloroethane	28 (35-120)	-	32 (≤25)		
	1,2,4-Trichlorobenzene	41 (45-120)	-	35 (≤20)		
	Acenaphthene	-	-	33 (≤20)		
	Acenaphthylene	-	-	30 (≤20)		
	Anthracene	-	-	23 (≤20)		
	Benzo(a)anthracene	-	-	24 (≤20)		
	Benzo(k)fluoranthene	-	-	23 (≤20)		
	Benzyl alcohol	-	-	33 (≤20)		
	Bis(2-chloroethoxy)methane	-	-	32 (≤20)		
	Bis(2-chloroethyl) ether	-	-	30 (≤20)		
	Bis(2-chloroisopropyl)ether	-	-	31 (≤20)		
	Bis(2-ethylhexyl)phthalate	-	-	25 (≤20)		
	4-Bromophenyl-phenyl ether	-	-	29 (≤25)		
	Butylbenzylphthalate	-	-	26 (≤20)		
	4-Chloroaniline	-	-	32 (≤25)		
	2-Chloronaphthalene	-	-	32 (≤20)		
	4-Chloro-3-methylphenol	-	-	33 (≤25)		
	2-Chlorophenol	-	-	30 (≤25)		
	4-Chlorophenyl-phenyl ether	-	-	33 (≤20)		
	Chrysene	-	-	28 (≤20)		
	Dibenzofuran	-	-	31 (≤20)		
	3,3'-Dichlorobenzidine	-	-	31 (≤25)		
	2,4-Dichlorophenol	-	-	33 (≤20)		
	2,4-Dimethylphenol	-	-	39 (≤25)		
	2,4-Dinitrophenol	-	-	26 (≤25)		
	2,4-Dinitrotoluene	-	-	24 (≤20)		
	2,6-Dinitrotoluene	-	-	30 (≤20)		
	Di-n-octylphthalate	-	-	30 (≤20)		
	Fluoranthene	-	-	21 (≤20)		
	Fluorene	-	-	34 (≤20)		
	Hexachlorobenzene	-	-	24 (≤20)		
	Hexachlorocyclopentadiene	-	-	70 (≤30)		
	Isophorone	-	-	32 (≤20)		
	2-Methylnaphthalene	-	-	31 (≤20)		
	2-Methylphenol	-	-	33 (≤20)		
	4-Methylphenol	-	-	30 (≤20)		
	Naphthalene	-	-	29 (≤20)		
2-Nitroaniline	-	-	32 (≤20)			
3-Nitroaniline	-	-	30 (≤25)			
4-Nitroaniline	-	-	27 (≤20)			
Nitrobenzene	-	-	34 (≤25)			
2-Nitrophenol	-	-	37 (≤25)			
N-Nitrosodiphenylamine	-	-	24 (≤20)			
N-Nitroso-di-n-propylamine	-	-	32 (≤20)			
Phenanthrene	-	-	22 (≤20)			
Phenol	-	-	29 (≤25)			
2,4,5-Trichlorophenol	-	-	35 (≤30)			
N-Nitrosodimethylamine	-	-	30 (≤20)			
1,2-Diphenylhydrazine/Azobenzene	-	-	27 (≤25)			

IX. Regional Quality Assurance and Quality Control

Not applicable.

X. Internal Standards

All internal standard areas and retention times were within QC limits.

XI. Target Compound Identifications

All target compound identifications were within validation criteria.

XII. Compound Quantitation and CRQLs

All compound quantitation and CRQLs were within validation criteria.

XIII. Tentatively Identified Compounds (TICs)

Tentatively identified compounds were not reported by the laboratory.

XIV. System Performance

The system performance was acceptable.

XV. Overall Assessment

Data flags have been summarized at the end of the report if data has been qualified.

XVI. Field Duplicates

No field duplicates were identified in this SDG.

XVII. Field Blanks

No field blanks were identified in this SDG.

**Boeing Realty Corp., Bldg C-1 Long Beach
Semivolatiles - Data Qualification Summary - SDG IQC1776**

SDG	Sample	Compound	Flag	A or P	Reason
IQC1776	MW3017_WG031507_0001	1,3-Dichlorobenzene 1,4-Dichlorobenzene 1,2-Dichlorobenzene Hexachlorobutadiene Hexachloroethane 1,2,4-Trichlorobenzene Acenaphthene Acenaphthylene Anthracene Benzo(a)anthracene Benzo(k)fluoranthene Benzyl alcohol Bis(2-chloroethoxy)methane Bis(2-chloroethyl) ether Bis(2-chloroisopropyl)ether Bis(2-ethylhexyl)phthalate 4-Bromophenyl-phenyl ether Butylbenzylphthalate 4-Chloroaniline 2-Chloronaphthalene 4-Chloro-3-methylphenol 2-Chlorophenol 4-Chlorophenyl-phenyl ether Chrysene Dibenzofuran 3,3'-Dichlorobenzidine 2,4-Dichlorophenol 2,4-Dimethylphenol 2,4-Dinitrophenol 2,4-Dinitrotoluene 2,6-Dinitrotoluene Di-n-octylphthalate Fluoranthene Fluorene Hexachlorobenzene Hexachlorocyclopentadiene Isophorone 2-Methylnaphthalene 2-Methylphenol 4-Methylphenol Naphthalene 2-Nitroaniline 3-Nitroaniline 4-Nitroaniline Nitrobenzene 2-Nitrophenol N-Nitrosodiphenylamine N-Nitroso-di-n-propylamine Phenanthrene Phenol 2,4,5-Trichlorophenol N-Nitrosodimethylamine 1,2-Diphenylhydrazine/Azobenzene	J (all detects) UJ (all non-detects)	P	Laboratory control samples (%R) (RPD)

**Boeing Realty Corp., Bldg C-1 Long Beach
Semivolatiles - Laboratory Blank Data Qualification Summary - SDG IQC1776**

No Sample Data Qualified in this SDG

TestAmerica

ANALYTICAL TESTING CORPORATION

17461 Derian Avenue, Suite 100, Irvine, CA 92614 (949) 261-1022 Fax: (949) 260-3297

TAIT Environmental/Boeing 701 N. Parkcenter Drive Santa Ana, CA 92705 Attention: Mehmet Pehlivan	Project ID: Boeing C-1 Long Beach EM-2701 Report Number: IQC1776	Sampled: 03/15/07 Received: 03/15/07
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SEMI-VOLATILE ORGANICS BY GC/MS (EPA 3520C/8270C)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC1776-04 (MW3017_WG031507_0001 - Water)									
Reporting Units: ug/l									
Acenaphthene	EPA 8270C	7C16066	1.9	9.5	ND UJ	0.948	03/16/07	03/20/07	
Acenaphthylene	EPA 8270C	7C16066	1.9	9.5	ND ↓	0.948	03/16/07	03/20/07	
Aniline	EPA 8270C	7C16066	2.4	9.5	ND	0.948	03/16/07	03/20/07	
Anthracene	EPA 8270C	7C16066	1.9	9.5	ND UJ	0.948	03/16/07	03/20/07	
Benzidine	EPA 8270C	7C16066	8.1	19	ND	0.948	03/16/07	03/20/07	
Benzoic acid	EPA 8270C	7C16066	8.1	19	ND	0.948	03/16/07	03/20/07	
Benzo(a)anthracene	EPA 8270C	7C16066	1.9	9.5	ND UJ	0.948	03/16/07	03/20/07	
Benzo(b)fluoranthene	EPA 8270C	7C16066	1.9	9.5	ND	0.948	03/16/07	03/20/07	
Benzo(k)fluoranthene	EPA 8270C	7C16066	1.9	9.5	ND UJ	0.948	03/16/07	03/20/07	
Benzo(g,h,i)perylene	EPA 8270C	7C16066	2.8	9.5	ND	0.948	03/16/07	03/20/07	
Benzo(a)pyrene	EPA 8270C	7C16066	1.9	9.5	ND	0.948	03/16/07	03/20/07	
Benzyl alcohol	EPA 8270C	7C16066	2.4	19	ND UJ	0.948	03/16/07	03/20/07	
Bis(2-chloroethoxy)methane	EPA 8270C	7C16066	1.9	9.5	ND	0.948	03/16/07	03/20/07	
Bis(2-chloroethyl)ether	EPA 8270C	7C16066	2.4	9.5	ND	0.948	03/16/07	03/20/07	
Bis(2-chloroisopropyl)ether	EPA 8270C	7C16066	2.4	9.5	ND	0.948	03/16/07	03/20/07	
Bis(2-ethylhexyl)phthalate	EPA 8270C	7C16066	3.8	47	ND	0.948	03/16/07	03/20/07	
4-Bromophenyl phenyl ether	EPA 8270C	7C16066	2.4	9.5	ND	0.948	03/16/07	03/20/07	
Butyl benzyl phthalate	EPA 8270C	7C16066	3.8	19	ND	0.948	03/16/07	03/20/07	
4-Chloroaniline	EPA 8270C	7C16066	1.9	9.5	ND	0.948	03/16/07	03/20/07	
2-Chloronaphthalene	EPA 8270C	7C16066	1.9	9.5	ND	0.948	03/16/07	03/20/07	
4-Chloro-3-methylphenol	EPA 8270C	7C16066	1.9	19	ND	0.948	03/16/07	03/20/07	
2-Chlorophenol	EPA 8270C	7C16066	1.9	9.5	ND	0.948	03/16/07	03/20/07	
4-Chlorophenyl phenyl ether	EPA 8270C	7C16066	1.9	9.5	ND	0.948	03/16/07	03/20/07	
Chrysene	EPA 8270C	7C16066	1.9	9.5	ND ✓	0.948	03/16/07	03/20/07	
Dibenz(a,h)anthracene	EPA 8270C	7C16066	2.8	19	ND	0.948	03/16/07	03/20/07	
Dibenzofuran	EPA 8270C	7C16066	1.9	9.5	ND UJ	0.948	03/16/07	03/20/07	
Di-n-butyl phthalate	EPA 8270C	7C16066	1.9	19	ND	0.948	03/16/07	03/20/07	
1,3-Dichlorobenzene	EPA 8270C	7C16066	2.8	9.5	ND UJ	0.948	03/16/07	03/20/07	L2
1,4-Dichlorobenzene	EPA 8270C	7C16066	2.4	9.5	ND ↓	0.948	03/16/07	03/20/07	L2
1,2-Dichlorobenzene	EPA 8270C	7C16066	2.8	9.5	ND ↓	0.948	03/16/07	03/20/07	L2
3,3-Dichlorobenzidine	EPA 8270C	7C16066	2.8	19	ND	0.948	03/16/07	03/20/07	
2,4-Dichlorophenol	EPA 8270C	7C16066	1.9	9.5	ND ✓	0.948	03/16/07	03/20/07	
Diethyl phthalate	EPA 8270C	7C16066	1.9	9.5	ND	0.948	03/16/07	03/20/07	
2,4-Dimethylphenol	EPA 8270C	7C16066	3.3	19	ND UJ	0.948	03/16/07	03/20/07	
Dimethyl phthalate	EPA 8270C	7C16066	1.9	9.5	ND	0.948	03/16/07	03/20/07	
4,6-Dinitro-2-methylphenol	EPA 8270C	7C16066	3.8	19	ND	0.948	03/16/07	03/20/07	
2,4-Dinitrophenol	EPA 8270C	7C16066	4.3	19	ND UJ	0.948	03/16/07	03/20/07	
2,4-Dinitrotoluene	EPA 8270C	7C16066	1.9	9.5	ND ↓	0.948	03/16/07	03/20/07	
2,6-Dinitrotoluene	EPA 8270C	7C16066	1.9	9.5	ND	0.948	03/16/07	03/20/07	
Di-n-octyl phthalate	EPA 8270C	7C16066	1.9	19	ND	0.948	03/16/07	03/20/07	
Fluoranthene	EPA 8270C	7C16066	1.9	9.5	ND ↓	0.948	03/16/07	03/20/07	

TestAmerica - Irvine, CA
Nicholas Marz
Project Manager

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TestAmerica

ANALYTICAL TESTING CORPORATION

17461 Derian Avenue, Suite 100, Irvine, CA 92614 (949) 261-1022 Fax: (949) 260-3297

TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-1 Long Beach
EM-2701
Report Number: IQC1776

Sampled: 03/15/07
Received: 03/15/07

SEMI-VOLATILE ORGANICS BY GC/MS (EPA 3520C/8270C)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC1776-04 (MW3017_WG031507_0001 - Water) - cont.									
Reporting Units: ug/l									
Fluorene	EPA 8270C	7C16066	1.9	9.5	ND	0.948	03/16/07	03/20/07	
Hexachlorobenzene	EPA 8270C	7C16066	2.4	9.5	ND	0.948	03/16/07	03/20/07	
Hexachlorobutadiene	EPA 8270C	7C16066	3.3	9.5	ND	0.948	03/16/07	03/20/07	L2
Hexachlorocyclopentadiene	EPA 8270C	7C16066	4.7	19	ND	0.948	03/16/07	03/20/07	
Hexachloroethane	EPA 8270C	7C16066	2.8	9.5	ND	0.948	03/16/07	03/20/07	L2
Indeno(1,2,3-cd)pyrene	EPA 8270C	7C16066	2.8	19	ND	0.948	03/16/07	03/20/07	
Isophorone	EPA 8270C	7C16066	1.9	9.5	ND	0.948	03/16/07	03/20/07	
2-Methylnaphthalene	EPA 8270C	7C16066	1.9	9.5	13	0.948	03/16/07	03/20/07	
2-Methylphenol	EPA 8270C	7C16066	1.9	9.5	ND	0.948	03/16/07	03/20/07	
4-Methylphenol	EPA 8270C	7C16066	1.9	9.5	ND	0.948	03/16/07	03/20/07	
Naphthalene	EPA 8270C	7C16066	2.4	9.5	16	0.948	03/16/07	03/20/07	
2-Nitroaniline	EPA 8270C	7C16066	1.9	19	ND	0.948	03/16/07	03/20/07	
3-Nitroaniline	EPA 8270C	7C16066	1.9	19	ND	0.948	03/16/07	03/20/07	
4-Nitroaniline	EPA 8270C	7C16066	2.4	19	ND	0.948	03/16/07	03/20/07	
Nitrobenzene	EPA 8270C	7C16066	2.4	19	ND	0.948	03/16/07	03/20/07	
2-Nitrophenol	EPA 8270C	7C16066	3.3	9.5	ND	0.948	03/16/07	03/20/07	
4-Nitrophenol	EPA 8270C	7C16066	5.2	19	ND	0.948	03/16/07	03/20/07	
N-Nitrosodiphenylamine	EPA 8270C	7C16066	1.9	9.5	ND	0.948	03/16/07	03/20/07	
N-Nitroso-di-n-propylamine	EPA 8270C	7C16066	2.4	9.5	ND	0.948	03/16/07	03/20/07	C
Pentachlorophenol	EPA 8270C	7C16066	3.3	19	ND	0.948	03/16/07	03/20/07	
Phenanthrene	EPA 8270C	7C16066	1.9	9.5	ND	0.948	03/16/07	03/20/07	
Phenol	EPA 8270C	7C16066	1.9	9.5	ND	0.948	03/16/07	03/20/07	
Pyrene	EPA 8270C	7C16066	1.9	9.5	ND	0.948	03/16/07	03/20/07	
1,2,4-Trichlorobenzene	EPA 8270C	7C16066	2.4	9.5	ND	0.948	03/16/07	03/20/07	L2
2,4,5-Trichlorophenol	EPA 8270C	7C16066	2.8	19	ND	0.948	03/16/07	03/20/07	
2,4,6-Trichlorophenol	EPA 8270C	7C16066	2.8	19	ND	0.948	03/16/07	03/20/07	
N-Nitrosodimethylamine	EPA 8270C	7C16066	2.4	19	ND	0.948	03/16/07	03/20/07	
1,2-Diphenylhydrazine/Azobenzene	EPA 8270C	7C16066	1.9	19	ND	0.948	03/16/07	03/20/07	C
Surrogate: 2-Fluorophenol (30-120%)					65 %				
Surrogate: Phenol-d6 (35-120%)					73 %				
Surrogate: 2,4,6-Tribromophenol (40-120%)					83 %				
Surrogate: Nitrobenzene-d5 (40-120%)					79 %				
Surrogate: 2-Fluorobiphenyl (45-120%)					67 %				
Surrogate: Terphenyl-d14 (45-120%)					71 %				

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Nicholas Marz
Project Manager

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LDC #: 16591B2
 SDG #: IQC1776
 Laboratory: Test America

VALIDATION COMPLETENESS WORKSHEET
 Tier 3

Date: 4/26/07
 Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: GC/MS Semivolatiles (EPA SW 846 Method 8270C)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: <u>3/5/07</u>
II.	GC/MS Instrument performance check	A	
III.	Initial calibration	A	% PSD, $r^2 = 0.990$
IV.	Continuing calibration	A	ICV = <u>25</u>
V.	Blanks	A	
VI.	Surrogate spikes	A	
VII.	Matrix spike/Matrix spike duplicates	N	client specified
VIII.	Laboratory control samples	SW	see IP
IX.	Regional Quality Assurance and Quality Control	N	
X.	Internal standards	A	
XI.	Target compound identification	A	
XII.	Compound quantitation/CRQLs	A	
XIII.	Tentatively identified compounds (TICs)	N	not reported
XIV.	System performance	A	
XV.	Overall assessment of data	A	
XVI.	Field duplicates	N	
XVII.	Field blanks	N	

Note: A = Acceptable ND = No compounds detected D = Duplicate
 N = Not provided/applicable R = Rinsate TB = Trip blank
 SW = See worksheet FB = Field blank EB = Equipment blank

Validated Samples: water

1	MW3017-WG031507_0001	11	7C16066-BLK	21		31	
2		12		22		32	
3		13		23		33	
4		14		24		34	
5		15		25		35	
6		16		26		36	
7		17		27		37	
8		18		28		38	
9		19		29		39	
10		20		30		40	

LDC #: 16591B2
 SDG #: see cover

VALIDATION FINDINGS CHECKLIST

Page: 1 of 2
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

Method: Semivolatiles (EPA SW 846 Method 8270C)

Validation Area	Yes	No	NA	Findings/Comments
All technical holding times were met.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Cooler temperature criteria was met.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the DFTPP performance results reviewed and found to be within the specified criteria?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all samples analyzed within the 12 hour clock criteria?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Did the laboratory perform a 5 point calibration prior to sample analysis?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all percent relative standard deviations (%RSD) and relative response factors (RRF) within method criteria for all CCCs and SPCCs?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a curve fit used for evaluation?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Did the initial calibration meet the curve fit acceptance criteria of ≥ 0.990 ?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all percent relative standard deviations (%RSD) $\leq 30\%$ and relative response factors (RRF) > 0.05 ?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a continuing calibration standard analyzed at least once every 12 hours for each instrument?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all percent differences (%D) and relative response factors (RRF) within method criteria for all CCCs and SPCCs?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all percent differences (%D) $\leq 25\%$ and relative response factors (RRF) ≥ 0.05 ?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a method blank associated with every sample in this SDG?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a method blank analyzed for each matrix and concentration?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was there contamination in the method blanks? If yes, please see the Blanks validation completeness worksheet.	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were all surrogate %R within QC limits?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
If 2 or more base neutral or acid surrogates were outside QC limits, was a reanalysis performed to confirm %R?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
If any %R was less than 10 percent, was a reanalysis performed to confirm %R?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were a matrix spike (MS) and matrix spike duplicate (MSD) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD. Soil / Water.	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Was a MS/MSD analyzed every 20 samples of each matrix?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the QC limits?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Was an LCS analyzed for this SDG?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	

LDC #: 16591B2
 SDG #: per cover

VALIDATION FINDINGS CHECKLIST

Page: 2 of 2
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

Validation Area	Yes	No	NA	Findings/Comments
Was an LCS analyzed per extraction batch?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the QC limits?	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
Were performance evaluation (PE) samples performed?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were the performance evaluation (PE) samples within the acceptance limits?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were internal standard area counts within -50% or +100% of the associated calibration standard?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were retention times within + 30 seconds from the associated calibration standard?	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were relative retention times (RRT's) within + 0.06 RRT units of the standard?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Did compound spectra meet specified EPA "Functional Guidelines" criteria?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were chromatogram peaks verified and accounted for?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the correct internal standard (IS), quantitation ion and relative response factor (RRF) used to quantitate the compound?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were compound quantitation and CRQLs adjusted to reflect all sample dilutions and dry weight factors applicable to level IV validation?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the major ions (> 10 percent relative intensity) in the reference spectrum evaluated in sample spectrum?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were relative intensities of the major ions within $\pm 20\%$ between the sample and the reference spectra?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Did the raw data indicate that the laboratory performed a library search for all required peaks in the chromatograms (samples and blanks)?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
System performance was found to be acceptable.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Overall assessment of data was found to be acceptable.	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Field duplicate pairs were identified in this SDG.	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
Target compounds were detected in the field duplicates.	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Field blanks were identified in this SDG.	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
Target compounds were detected in the field blanks.	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	

VALIDATION FINDINGS WORKSHEET

METHOD: GC/MS BNA (EPA SW 846 Method 8270)

A. Phenol**	P. Bis(2-chloroethoxy)methane	EE. 2,6-Dinitrotoluene	TT. Pentachlorophenol**	III. Benzo(a)pyrene**
B. Bis (2-chloroethyl) ether	Q. 2,4-Dichlorophenol**	FF. 3-Nitroaniline	UU. Phenanthrene	JJJ. Indeno(1,2,3-cd)pyrene
C. 2-Chlorophenol	R. 1,2,4-Trichlorobenzene	GG. Acenaphthene**	VV. Anthracene	KKK. Dibenz(a,h)anthracene
D. 1,3-Dichlorobenzene	S. Naphthalene	HH. 2,4-Dinitrophenol*	WW. Carbazole	LLL. Benzo(g,h,i)perylene
E. 1,4-Dichlorobenzene**	T. 4-Chloroaniline	II. 4-Nitrophenol*	XX. Di-n-butylphthalate	MMM. Bis(2-Chloroisopropyl)ether
F. 1,2-Dichlorobenzene	U. Hexachlorobutadiene**	JJ. Dibenzofuran	YY. Fluoranthene**	NNN. Aniline
G. 2-Methylphenol	V. 4-Chloro-3-methylphenol**	KK. 2,4-Dinitrotoluene	ZZ. Pyrene	OOO. N-Nitrosodimethylamine
H. 2,2'-Oxybis(1-chloropropane)	W. 2-Methylnaphthalene	LL. Diethylphthalate	AAA. Butylbenzylphthalate	PPP. Benzoic Acid
I. 4-Methylphenol	X. Hexachlorocyclopentadiene*	MM. 4-Chlorophenyl-phenyl ether	BBB. 3,3'-Dichlorobenzidine	QQQ. Benzyl alcohol
J. N-Nitroso-di-n-propylamine*	Y. 2,4,6-Trichlorophenol**	NN. Fluorene	CCC. Benzo(a)anthracene	RRR. Pyridine
K. Hexachloroethane	Z. 2,4,5-Trichlorophenol	OO. 4-Nitroaniline	DDD. Chrysene	SSS. Benzidine
L. Nitrobenzene	AA. 2-Chloronaphthalene	PP. 4,6-Dinitro-2-methylphenol	EEE. Bis(2-ethylhexyl)phthalate	TTT. <i>Azo benzene</i>
M. Isophorone	BB. 2-Nitroaniline	QQ. N-Nitrosodiphenylamine (1)**	FFF. Di-n-octylphthalate**	UUU.
N. 2-Nitrophenol**	CC. Dimethylphthalate	RR. 4-Bromophenyl-phenylether	GGG. Benzo(b)fluoranthene	VVV.
O. 2,4-Dimethylphenol	DD. Acenaphthylene	SS. Hexachlorobenzene	HHH. Benzo(k)fluoranthene	WWW.

COMPNDL2S

TAIT Environmental/Boeing 701 N. Parkcenter Drive Santa Ana, CA 92705 Attention: Mehmet Pehlivan	Project ID: Boeing C-1 Long Beach EM-2701 Report Number: IQC1776	Sampled: 03/15/07 Received: 03/15/07
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METHOD BLANK/OC DATA

SEMI-VOLATILE ORGANICS BY GC/MS (EPA 3520C/8270C)

Analyte	Result	Reporting Limit	MDL	Units	Spike Level	Source Result	%REC	%REC Limits	RPD	RPD Limit	Data Qualifiers
Batch: 7C16066 Extracted: 03/16/07											
Blank Analyzed: 03/19/2007 (7C16066-BLK1)											
Surrogate: Phenol-d6	13.6			ug/l	20.0		68	35-120			
Surrogate: 2,4,6-Tribromophenol	13.2			ug/l	20.0		66	40-120			
Surrogate: Nitrobenzene-d5	5.90			ug/l	10.0		59	40-120			
Surrogate: 2-Fluorobiphenyl	7.14			ug/l	10.0		71	45-120			
Surrogate: Terphenyl-d14	7.38			ug/l	10.0		74	45-120			
LCS Analyzed: 03/19/2007 (7C16066-BS1)											
Acenaphthene	64.6	10	2.0	ug/l	100		65	55-120			MNR1
Acenaphthylene	72.4	10	2.0	ug/l	100		72	60-120			
Aniline	78.3	10	2.5	ug/l	100		78	40-120			
Anthracene	72.1	10	2.0	ug/l	100		72	60-120			
Benzidine	147	20	8.5	ug/l	100		147	25-160			
Benzoic acid	31.7	20	8.5	ug/l	100		32	25-120			
Benzo(a)anthracene	71.5	10	2.0	ug/l	100		72	60-120			
Benzo(b)fluoranthene	82.5	10	2.0	ug/l	100		82	55-125			
Benzo(k)fluoranthene	82.9	10	2.0	ug/l	100		83	50-125			
Benzo(g,h,i)perylene	98.0	10	3.0	ug/l	100		98	45-130			
Benzo(a)pyrene	88.8	10	2.0	ug/l	100		89	55-125			
Benzyl alcohol	61.3	20	2.5	ug/l	100		61	50-120			
Bis(2-chloroethoxy)methane	62.3	10	2.0	ug/l	100		62	55-120			
Bis(2-chloroethyl)ether	54.6	10	2.5	ug/l	100		55	50-120			
Bis(2-chloroisopropyl)ether	55.2	10	2.5	ug/l	100		55	45-120			
Bis(2-ethylhexyl)phthalate	69.2	50	4.0	ug/l	100		69	60-125			
4-Bromophenyl phenyl ether	66.5	10	2.5	ug/l	100		66	55-120			
Butyl benzyl phthalate	68.1	20	4.0	ug/l	100		68	50-125			
4-Chloroaniline	63.7	10	2.0	ug/l	100		64	50-120			
2-Chloronaphthalene	61.8	10	2.0	ug/l	100		62	55-120			
4-Chloro-3-methylphenol	61.4	20	2.0	ug/l	100		61	55-120			
2-Chlorophenol	57.6	10	2.0	ug/l	100		58	45-120			
4-Chlorophenyl phenyl ether	63.9	10	2.0	ug/l	100		64	60-120			
Chrysene	69.5	10	2.0	ug/l	100		70	60-120			
Dibenz(a,h)anthracene	94.5	20	3.0	ug/l	100		94	50-135			
Dibenzofuran	64.0	10	2.0	ug/l	100		64	60-120			
Di-n-butyl phthalate	75.7	20	2.0	ug/l	100		76	55-125			
1,3-Dichlorobenzene	29.3	10	3.0	ug/l	100		29	35-120			L2
1,4-Dichlorobenzene	34.4	10	2.5	ug/l	100		34	35-120			L2

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Project Manager

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TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-1 Long Beach
EM-2701
Report Number: IQC1776

Sampled: 03/15/07
Received: 03/15/07



SEMI-VOLATILE ORGANICS BY GC/MS (EPA 3520C/8270C)

Analyte	Result	Reporting Limit	MDL	Units	Spike Level	Source Result	%REC	%REC Limits	RPD	RPD Limit	Data Qualifiers
Batch: 7C16066 Extracted: 03/16/07											
LCS Analyzed: 03/19/2007 (7C16066-BS1)											
1,2-Dichlorobenzene	35.3	10	3.0	ug/l	100		35	40-120	F		MNR1
3,3-Dichlorobenzidine	68.8	20	3.0	ug/l	100		69	50-135			L2
2,4-Dichlorophenol	56.7	10	2.0	ug/l	100		57	50-120			
Diethyl phthalate	69.3	10	2.0	ug/l	100		69	50-120			
2,4-Dimethylphenol	45.8	20	3.5	ug/l	100		46	35-120			
Dimethyl phthalate	60.9	10	2.0	ug/l	100		61	25-120			
4,6-Dinitro-2-methylphenol	71.9	20	4.0	ug/l	100		72	40-120			
2,4-Dinitrophenol	68.7	20	4.5	ug/l	100		69	35-120			
2,4-Dinitrotoluene	74.4	10	2.0	ug/l	100		74	60-120			
2,6-Dinitrotoluene	67.9	10	2.0	ug/l	100		68	60-120			
Di-n-octyl phthalate	70.1	20	2.0	ug/l	100		70	60-130			
Fluoranthene	76.3	10	2.0	ug/l	100		76	55-120			
Fluorene	63.5	10	2.0	ug/l	100		64	60-120			
Hexachlorobenzene	69.1	10	2.5	ug/l	100		69	55-120			
Hexachlorobutadiene	36.9	10	3.5	ug/l	100		37	40-120	U		L2
Hexachlorocyclopentadiene	34.2	20	5.0	ug/l	100		34	20-120			
Hexachloroethane	27.9	10	3.0	ug/l	100		28	35-120	K		L2
Indeno(1,2,3-cd)pyrene	95.4	20	3.0	ug/l	100		95	45-135			
Isophorone	52.4	10	2.0	ug/l	100		52	50-120			
2-Methylnaphthalene	58.0	10	2.0	ug/l	100		58	50-120			
2-Methylphenol	59.6	10	2.0	ug/l	100		60	50-120			
4-Methylphenol	63.4	10	2.0	ug/l	100		63	45-120			
Naphthalene	55.7	10	2.5	ug/l	100		56	50-120			
2-Nitroaniline	66.6	20	2.0	ug/l	100		67	60-120			
3-Nitroaniline	82.9	20	2.0	ug/l	100		83	55-120			
4-Nitroaniline	85.9	20	2.5	ug/l	100		86	50-125			
Nitrobenzene	52.0	20	2.5	ug/l	100		52	50-120			
2-Nitrophenol	58.6	10	3.5	ug/l	100		59	45-120			
4-Nitrophenol	68.6	20	5.5	ug/l	100		69	40-120			
N-Nitrosodiphenylamine	64.2	10	2.0	ug/l	100		64	55-120			
N-Nitroso-di-n-propylamine	54.1	10	2.5	ug/l	100		54	45-120			
Pentachlorophenol	83.2	20	3.5	ug/l	100		83	45-125			
Phenanthrene	70.3	10	2.0	ug/l	100		70	60-120			
Phenol	58.2	10	2.0	ug/l	100		58	45-120			
Pyrene	67.8	10	2.0	ug/l	100		68	50-125			

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Nicholas Marz
Project Manager

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TAIT Environmental/Boeing 701 N. Parkcenter Drive Santa Ana, CA 92705 Attention: Mehmet Pehlivan	Project ID: Boeing C-1 Long Beach EM-2701 Report Number: IQC1776	Sampled: 03/15/07 Received: 03/15/07
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METHOD BLANK/QC DATA

SEMI-VOLATILE ORGANICS BY GC/MS (EPA 3520C/8270C)

Analyte	Result	Reporting Limit	MDL	Units	Spike Level	Source Result	%REC %REC	Limits	RPD	RPD Limit	Data Qualifiers
Batch: 7C16066 Extracted: 03/16/07											
LCS Analyzed: 03/19/2007 (7C16066-BS1)											
1,2,4-Trichlorobenzene	40.7	10	2.5	ug/l	100		41	45-120	R		MNRI L2
2,4,5-Trichlorophenol	61.9	20	3.0	ug/l	100		62	50-120			
2,4,6-Trichlorophenol	63.2	20	3.0	ug/l	100		63	50-120			
N-Nitrosodimethylamine	51.3	20	2.5	ug/l	100		51	40-120			
1,2-Diphenylhydrazine/Azobenzene	64.0	20	2.0	ug/l	100		64	55-120			
Surrogate: 2-Fluorophenol	10.9			ug/l	20.0		54	30-120			
Surrogate: Phenol-d6	11.4			ug/l	20.0		57	35-120			
Surrogate: 2,4,6-Tribromophenol	14.1			ug/l	20.0		70	40-120			
Surrogate: Nitrobenzene-d5	5.54			ug/l	10.0		55	40-120			
Surrogate: 2-Fluorobiphenyl	6.42			ug/l	10.0		64	45-120			
Surrogate: Terphenyl-d14	6.96			ug/l	10.0		70	45-120			
LCS Dup Analyzed: 03/19/2007 (7C16066-BSD1)											
Acenaphthene	90.2	10	2.0	ug/l	100		90	55-120	33	20	GG R-7
Acenaphthylene	98.1	10	2.0	ug/l	100		98	60-120	30	20	DD R-7
Aniline	82.9	10	2.5	ug/l	100		83	40-120	6	30	
Anthracene	90.8	10	2.0	ug/l	100		91	60-120	23	20	WW R-7
Benzidine	149	20	8.5	ug/l	100		149	25-160	1	35	
Benzoic acid	32.5	20	8.5	ug/l	100		32	25-120	2	30	
Benzo(a)anthracene	90.6	10	2.0	ug/l	100		91	60-120	24	20	CCC R-7
Benzo(b)fluoranthene	99.2	10	2.0	ug/l	100		99	55-125	18	25	
Benzo(k)fluoranthene	104	10	2.0	ug/l	100		104	50-125	23	20	HHH R-7
Benzo(g,h,i)perylene	117	10	3.0	ug/l	100		117	45-130	18	25	
Benzo(a)pyrene	108	10	2.0	ug/l	100		108	55-125	20	25	
Benzyl alcohol	85.6	20	2.5	ug/l	100		86	50-120	33	20	QQQ R-7
Bis(2-chloroethoxy)methane	85.8	10	2.0	ug/l	100		86	55-120	32	20	P R-7
Bis(2-chloroethyl)ether	74.2	10	2.5	ug/l	100		74	50-120	30	20	B R-7
Bis(2-chloroisopropyl)ether	75.8	10	2.5	ug/l	100		76	45-120	31	20	MMM R-7
Bis(2-ethylhexyl)phthalate	89.2	50	4.0	ug/l	100		89	60-125	25	20	EEE R-7
4-Bromophenyl phenyl ether	89.2	10	2.5	ug/l	100		89	55-120	29	25	RR R-7
Butyl benzyl phthalate	88.2	20	4.0	ug/l	100		88	50-125	26	20	AAA R-7
4-Chloroaniline	88.2	10	2.0	ug/l	100		88	50-120	32	25	T R-7
2-Chloronaphthalene	85.1	10	2.0	ug/l	100		85	55-120	32	20	AA R-7
4-Chloro-3-methylphenol	85.6	20	2.0	ug/l	100		86	55-120	33	25	V R-7
2-Chlorophenol	78.1	10	2.0	ug/l	100		78	45-120	30	25	C R-7
4-Chlorophenyl phenyl ether	89.0	10	2.0	ug/l	100		89	60-120	33	20	MM R-7

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ANALYTICAL TESTING CORPORATION

17461 Derian Avenue, Suite 100, Irvine, CA 92614 (949) 261-1022 Fax:(949) 260-3297

TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-1 Long Beach
EM-2701
Report Number: IQC1776

Sampled: 03/15/07
Received: 03/15/07

METHOD BLANK/QC DATA

SEMI-VOLATILE ORGANICS BY GC/MS (EPA 3520C/8270C)

Analyte	Result	Reporting Limit	MDL	Units	Spike Level	Source Result	%REC %REC	%REC Limits	RPD	RPD Limit	Data Qualifiers
Batch: 7C16066 Extracted: 03/16/07											
LCS Dup Analyzed: 03/19/2007 (7C16066-BSD1)											
Chrysene	92.3	10	2.0	ug/l	100	92	60-120	28	20	PVD	R-7
Dibenz(a,h)anthracene	113	20	3.0	ug/l	100	113	50-135	18	25		
Dibenzofuran	87.7	10	2.0	ug/l	100	88	60-120	31	20	JJ	R-7
Di-n-butyl phthalate	90.8	20	2.0	ug/l	100	91	55-125	18	20		
1,3-Dichlorobenzene	41.4	10	3.0	ug/l	100	41	35-120	34	25	P-	R-2
1,4-Dichlorobenzene	45.3	10	2.5	ug/l	100	45	35-120	27	25	E-	R-2
1,2-Dichlorobenzene	48.2	10	3.0	ug/l	100	48	40-120	31	25	F-	R-2
3,3-Dichlorobenzidine	93.8	20	3.0	ug/l	100	94	50-135	31	25	BBB	R-7
2,4-Dichlorophenol	79.1	10	2.0	ug/l	100	79	50-120	33	20	GG	R-7
Diethyl phthalate	88.0	10	2.0	ug/l	100	88	50-120	24	30		
2,4-Dimethylphenol	67.9	20	3.5	ug/l	100	68	35-120	39	25	Θ	R-7
Dimethyl phthalate	79.9	10	2.0	ug/l	100	80	25-120	27	30		
4,6-Dinitro-2-methylphenol	89.2	20	4.0	ug/l	100	89	40-120	21	25		
2,4-Dinitrophenol	89.6	20	4.5	ug/l	100	90	35-120	26	25	HH	R-7
2,4-Dinitrotoluene	94.7	10	2.0	ug/l	100	95	60-120	24	20	KK	R-7
2,6-Dinitrotoluene	91.8	10	2.0	ug/l	100	92	60-120	30	20	EE	R-7
Di-n-octyl phthalate	94.5	20	2.0	ug/l	100	94	60-130	30	20	FFF	R-7
Fluoranthene	94.1	10	2.0	ug/l	100	94	55-120	21	20	YY	R-7
Fluorene	89.1	10	2.0	ug/l	100	89	60-120	34	20	NN	R-7
Hexachlorobenzene	88.1	10	2.5	ug/l	100	88	55-120	24	20	SS	R-7
Hexachlorobutadiene	50.8	10	3.5	ug/l	100	51	40-120	32	25	U-	R-2
Hexachlorocyclopentadiene	71.2	20	5.0	ug/l	100	71	20-120	70	30	X	R-7
Hexachloroethane	38.6	10	3.0	ug/l	100	39	35-120	32	25	K-	R-2
Indeno(1,2,3-cd)pyrene	113	20	3.0	ug/l	100	113	45-135	17	25	HH	
Isophorone	72.0	10	2.0	ug/l	100	72	50-120	32	20	M	R-7
2-Methylnaphthalene	79.6	10	2.0	ug/l	100	80	50-120	31	20	W	R-7
2-Methylphenol	82.8	10	2.0	ug/l	100	83	50-120	33	20	G	R-7
4-Methylphenol	85.5	10	2.0	ug/l	100	86	45-120	30	20	I	R-7
Naphthalene	74.9	10	2.5	ug/l	100	75	50-120	29	20	S	R-7
2-Nitroaniline	92.3	20	2.0	ug/l	100	92	60-120	32	20	BB	R-7
3-Nitroaniline	112	20	2.0	ug/l	100	112	55-120	30	25	FF	R-7
4-Nitroaniline	113	20	2.5	ug/l	100	113	50-125	27	20	ΘΘ	R-7
Nitrobenzene	73.2	20	2.5	ug/l	100	73	50-120	34	25	L	R-7
2-Nitrophenol	85.2	10	3.5	ug/l	100	85	45-120	37	25	N	R-7
4-Nitrophenol	90.2	20	5.5	ug/l	100	90	40-120	X	30		

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Nicholas Marz
Project Manager

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TAIT Environmental/Boeing 701 N. Parkcenter Drive Santa Ana, CA 92705 Attention: Mehmet Pehlivan	Project ID: Boeing C-1 Long Beach EM-2701 Report Number: IQC1776	Sampled: 03/15/07 Received: 03/15/07
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METHOD BLANK/QC DATA

SEMI-VOLATILE ORGANICS BY GC/MS (EPA 3520C/8270C)

Analyte	Result	Reporting Limit	MDL	Units	Spike Level	Source Result	%REC %REC	Limits	RPD	RPD Limit	Data Qualifiers
Batch: 7C16066 Extracted: 03/16/07											
LCS Dup Analyzed: 03/19/2007 (7C16066-BS1)											
N-Nitrosodiphenylamine	81.3	10	2.0	ug/l	100	81	55-120	24	20	QQ	R-7
N-Nitroso-di-n-propylamine	74.7	10	2.5	ug/l	100	75	45-120	32	20	J	R-7
Pentachlorophenol	106	20	3.5	ug/l	100	106	45-125	24	25		
Phenanthrene	87.3	10	2.0	ug/l	100	87	60-120	22	20	UU	R-7
Phenol	78.3	10	2.0	ug/l	100	78	45-120	29	25	A	R-7
Pyrene	83.6	10	2.0	ug/l	100	84	50-125	21	25		
1,2,4-Trichlorobenzene	58.0	10	2.5	ug/l	100	58	45-120	35	20	R-	R-2
2,4,5-Trichlorophenol	88.2	20	3.0	ug/l	100	88	50-120	35	30	Z	R-7
2,4,6-Trichlorophenol	84.6	20	3.0	ug/l	100	85	50-120	29	30		
N-Nitrosodimethylamine	69.6	20	2.5	ug/l	100	70	40-120	30	20	ooo	R-7
1,2-Diphenylhydrazine/Azobenzene	84.3	20	2.0	ug/l	100	84	55-120	27	25		R-7
Surrogate: 2-Fluorophenol	14.9			ug/l	20.0	74	30-120				
Surrogate: Phenol-d6	15.9			ug/l	20.0	80	35-120				
Surrogate: 2,4,6-Tribromophenol	19.0			ug/l	20.0	95	40-120				
Surrogate: Nitrobenzene-d5	7.64			ug/l	10.0	76	40-120				
Surrogate: 2-Fluorobiphenyl	8.82			ug/l	10.0	88	45-120				
Surrogate: Terphenyl-d14	8.86			ug/l	10.0	89	45-120				

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VALIDATION FIN 03 WORKSHEET
Initial Calibration Calculation Verification

Page 1 of 1
Reviewer: [Signature]
2nd Reviewer: [Signature]

METHOD: GC/MS BNA (EPA SW 846 Method 8270)

The Relative Response Factor (RRF), average RRF, and percent relative standard deviation (%RSD) were recalculated for the compounds identified below using the following calculations:

$RRF = (A_x/C_x)/(A_s/C_s)$
 average RRF = sum of the RRFs/number of standards
 $%RSD = 100 * (S/X)$
 A_x = Area of compound, A_s = Area of associated internal standard
 C_x = Concentration of compound, C_s = Concentration of internal standard
 S = Standard deviation of the RRFs, X = Mean of the RRFs

#	Standard ID	Calibration Date	Compound (Reference Internal Standard)	Reported		Recalculated		Reported		Recalculated	
				RRF (std)	RRF (std)	RRF (std)	Average RRF (Initial)	Average RRF (Initial)	%RSD	%RSD	
1	KAL-8	3/3/07	Phenol (1st internal standard)	2.111	2.11	2.125	2.125	4.01	4.01	4.01	4.01
			Naphthalene (2nd internal standard)	1.036	1.036	1.030	1.030	5.90	5.90	5.90	5.90
			Fluorene (3rd internal standard)	1.321	1.321	1.268	1.268	9.44	9.44	9.44	9.44
			Benzo(a)anthracene Benzo(a)anthracene (4th internal standard)	1.093 1.093	1.093	1.059 1.059	1.059	7.16	7.16	7.16	7.16
			Bis(2-ethylhexyl)phthalate (5th internal standard)	1.396	1.396	1.308	1.308	3.67	3.67	3.67	3.67
			Benzo(a)pyrene (6th internal standard)	1.205	1.205	1.192	1.192	4.89	4.89	4.89	4.89
2			Phenol (1st internal standard)								
			Naphthalene (2nd internal standard)								
			Fluorene (3rd internal standard)								
			Pentachlorophenol (4th internal standard)								
			Bis(2-ethylhexyl)phthalate (5th internal standard)								
			Benzo(a)pyrene (6th internal standard)								
3			Phenol (1st internal standard)								
			Naphthalene (2nd internal standard)								
			Fluorene (3rd internal standard)								
			Pentachlorophenol (4th internal standard)								
			Bis(2-ethylhexyl)phthalate (5th internal standard)								
			Benzo(a)pyrene (6th internal standard)								

Comments: Refer to Initial Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 16591B2
 SDG #: ps co vel

VALIDATION FINDINGS WORKSHEET
Continuing Calibration Results Verification

Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: GC/MS BNA (EPA SW 846 Method 8270)

The percent difference (%D) of the initial calibration average Relative Response Factors (RRFs) and the continuing calibration RRFs were recalculated for the compounds identified below using the following calculation:

% Difference = $100 * (\text{ave. RRF} - \text{RRF}) / \text{ave. RRF}$ Where: ave. RRF = initial calibration average RRF
 RRF = $(A_x)(C_s) / (A_s)(C_x)$ RRF = continuing calibration RRF
 A_x = Area of compound, A_s = Area of associated internal standard
 C_x = Concentration of compound, C_s = Concentration of internal standard

#	Standard ID	Calibration Date	Compound (Reference Internal Standard)	Average RRF (Initial)	Reported		Recalculated	
					RRF (CC)	%D	RRF (CC)	%D
1	cen	3/20/07	Phenol (1st internal standard)	2.125	2.072	2.5	2.072	2.5
			Naphthalene (2nd internal standard)	1.030	1.092	6.0	1.092	6.0
			Fluorene (3rd internal standard)	1.268	1.323	4.3	1.323	4.3
			Pentachlorophenol (4th internal standard)	1.089	1.117	5.5	1.117	5.5
			Bis(2-ethylhexyl)phthalate (5th internal standard)	1.308	1.392	6.4	1.392	6.4
			Benzo(e)pyrene (6th internal standard)	1.192	1.192	0.0	1.192	0.0
2			Phenol (1st internal standard)					
			Naphthalene (2nd internal standard)					
			Fluorene (3rd internal standard)					
			Pentachlorophenol (4th internal standard)					
			Bis(2-ethylhexyl)phthalate (5th internal standard)					
			Benzo(e)pyrene (6th internal standard)					
3			Phenol (1st internal standard)					
			Naphthalene (2nd internal standard)					
			Fluorene (3rd internal standard)					
			Pentachlorophenol (4th internal standard)					
			Bis(2-ethylhexyl)phthalate (5th internal standard)					
			Benzo(e)pyrene (6th internal standard)					

Comments: Refer to Continuing Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 16591B2
 SDG #: see cover

VALIDATION FINDINGS WORKSHEET
Surrogate Results Verification

Page: 1 of 1
 Reviewer: A
 2nd reviewer: N

METHOD: GC/MS Semivolatiles (EPA SW 846 Method 8270)

The percent recoveries (%R) of surrogates were recalculated for the compounds identified below using the following calculation:

% Recovery: $SF/SS * 100$

Where: SF = Surrogate Found
 SS = Surrogate Spiked

Sample ID: # 1

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Nitrobenzene-d5	5	3.95	79	79	0
2-Fluorobiphenyl	↓	3.37	67	67	↓
Terphenyl-d14	↓	3.56	71	71	↓
Phenol-d5	10	7.29	73	73	↓
2-Fluorophenol	↓	6.49	65	65	↓
2,4,6-Tribromophenol	↓	8.30	83	83	↓
2-Chlorophenol-d4					
1,2-Dichlorobenzene-d4					

Sample ID: _____

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Nitrobenzene-d5					
2-Fluorobiphenyl					
Terphenyl-d14					
Phenol-d5					
2-Fluorophenol					
2,4,6-Tribromophenol					
2-Chlorophenol-d4					
1,2-Dichlorobenzene-d4					

Sample ID: _____

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Nitrobenzene-d5					
2-Fluorobiphenyl					
Terphenyl-d14					
Phenol-d5					
2-Fluorophenol					
2,4,6-Tribromophenol					
2-Chlorophenol-d4					
1,2-Dichlorobenzene-d4					

LDC #: 16591B2
 SDG #: 16591B2

VALIDATION FINDINGS WORKSHEET
Laboratory Control Sample/Laboratory Control Sample Duplicates Results Verification

Page: 1 of 7
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: GC/MS BNA (EPA SW 846 Method 8270)

The percent recoveries (%R) and Relative Percent Difference (RPD) of the laboratory control sample and laboratory control sample duplicate were recalculated for the compounds identified below using the following calculation:

% Recovery = $100 * (SC/SA)$

Where: SSC = Spike concentration
 SA = Spike added

RPD = $100 * (LCS - LCSD) / (LCS + LCSD)$

LCS = Laboratory control sample percent recovery LCSD = Laboratory control sample duplicate percent recovery

LCS/LCSD samples: 7016066 - LCS 1D

Compound	Spike Added (ng/L)		Spike Concentration (ng/L)		LCS		LCSD		LCS		LCSD	
	LCS	LCSD	LCS	LCSD	Reported	Recalc.	Reported	Recalc.	Reported	Recalc.	Reported	Recalc.
	Percent Recovery		Percent Recovery		Percent Recovery		Percent Recovery		Percent Recovery		Percent Recovery	
Phenol	100		58.2	78.3	58	50	78	78	29	29	29	29
N-Nitroso-dl-n-propylamine			54.1	74.7	54	54	75	75	32	32	32	32
4-Chloro-3-methylphenol			61.4	85.6	61	61	86	86	33	33	33	33
Acenaphthene			64.6	90.2	65	65	90	90	33	33	33	33
Pentachlorophenol			83.2	106	83	83	106	106	24	24	24	24
Pyrene			67.8	83.6	68	60	84	84	21	21	21	21

Comments: Refer to Laboratory Control Sample/Laboratory Control Sample Duplicates findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

**Boeing Realty Corp., Bldg C-1 Long Beach
Data Validation Reports
LDC# 16591**

Hexavalent Chromium

LDC

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Boeing Realty Corp., Bldg C-1 Long Beach
Collection Date: March 8, 2007
LDC Report Date: April 26, 2007
Matrix: Water
Parameters: Hexavalent chromium
Validation Level: Tier 1, 2, & 3
Laboratory: TestAmerica
Sample Delivery Group (SDG): IQC0980

Sample Identification

MW3009_WG030807_0001
MW3012_WG030807_0001*
MW3012_WG030807_0002**

*Indicates sample underwent Tier 2 review
**Indicates sample underwent Tier 3 review
All other samples underwent Tier 1 review

Introduction

This data review covers 3 water samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 7196A for Hexavalent chromium.

The review follows a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Inorganic Data Review (October 2004) as there are no current guidelines for the methods stated above.

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

Blank results are summarized in Section III.

Field duplicates are summarized in Section IX.

Samples indicated by a double asterisk on the front cover underwent a Tier 3 review. A Tier 2 or Tier 1 review was performed on all of the other samples. Raw data were not evaluated for the samples reviewed by Tier 2 or Tier 1 criteria since this review is based on QC data.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- N Presumptive evidence of presence of the constituent.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. Calibration

a. Initial Calibration

All criteria for the initial calibration of each method were met.

Initial calibration data were not reviewed for Tier I.

b. Calibration Verification

Calibration verification frequency and analysis criteria were met for each method when applicable.

Calibration verification data were not reviewed for Tier I.

III. Blanks

Method blanks were reviewed for each matrix as applicable. No contaminant concentrations were found in the initial, continuing and preparation blanks.

IV. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

V. Duplicates

Duplicate (DUP) sample analyses were reviewed for each matrix as applicable.

VI. Laboratory Control Samples

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) were within QC limits.

VII. Sample Result Verification

All sample result verifications were acceptable for samples on which a Tier 3 review was performed. Raw data were not evaluated for the samples reviewed by Tier 2 or Tier 1 criteria.

VIII. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

IX. Field Duplicates

Samples were identified as field duplicates. No contaminant concentrations were detected in any of the samples with the following exceptions:

Analyte	Concentration (ug/L)		RPD
	MW3012_WG030807_0001*	MW3012_WG030807_0002**	
Hexavalent chromium	1.3	0.98	28

X. Field Blanks

No field blanks were identified in this SDG.

**Boeing Realty Corp., Bldg C-1 Long Beach
Hexavalent chromium - Data Qualification Summary - SDG IQC0980**

No Sample Data Qualified in this SDG

**Boeing Realty Corp., Bldg C-1 Long Beach
Hexavalent chromium - Laboratory Blank Data Qualification Summary - SDG
IQC0980**

No Sample Data Qualified in this SDG

TestAmerica

ANALYTICAL TESTING CORPORATION

17461 Derian Avenue, Suite 100, Irvine, CA 92614 (949) 261-1022 Fax: (949) 260-3297

TAIT Environmental/Boeing
701 N Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-1 Long Beach
EM-2701-05
Report Number: IQC0980

Sampled: 03/08/07
Received: 03/08/07

INORGANICS

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC0980-01 (TB_TAIT030807_0001 - Water)					Sampled: 03/08/07				
Reporting Units: ug/l									
Chromium VI	EPA 7196A	7C08171	0.65	25	ND	1	03/08/07	03/08/07	
Sample ID: IQC0980-02 (MW3009_WG030807_0001 - Water)					Sampled: 03/08/07				
Reporting Units: ug/l									
Chromium VI	EPA 7196A	7C08171	0.65	25	ND	1	03/08/07	03/08/07	
Sample ID: IQC0980-03 (MW3016_WG030807_0001 - Water)					Sampled: 03/08/07				
Reporting Units: ug/l									
Chromium VI	EPA 7196A	7C08171	0.65	25	11	1	03/08/07	03/08/07	J
Sample ID: IQC0980-04 (MW3015_WG030807_0001 - Water)					Sampled: 03/08/07				
Reporting Units: ug/l									
Chromium VI	EPA 7196A	7C08171	0.65	25	49	1	03/08/07	03/08/07	
Sample ID: IQC0980-05 (MW3014_WG030807_0001 - Water)					Sampled: 03/08/07				
Reporting Units: ug/l									
Chromium VI	EPA 7196A	7C08171	0.65	25	13	1	03/08/07	03/08/07	J
Sample ID: IQC0980-06 (MW3013_WG030807_0001 - Water)					Sampled: 03/08/07				
Reporting Units: ug/l									
Chromium VI	EPA 7196A	7C08171	0.65	25	3.4	1	03/08/07	03/08/07	J
Sample ID: IQC0980-07 (MW3012_WG030807_0001 - Water)					Sampled: 03/08/07				
Reporting Units: ug/l									
Chromium VI	EPA 7196A	7C08171	0.65	25	1.3	1	03/08/07	03/08/07	J
Sample ID: IQC0980-08 (MW3012_WG030807_0002 - Water)					Sampled: 03/08/07				
Reporting Units: ug/l									
Chromium VI	EPA 7196A	7C08171	0.65	25	0.98	1	03/08/07	03/08/07	J
Sample ID: IQC0980-09 (MW3011_WG030807_0001 - Water)					Sampled: 03/08/07				
Reporting Units: ug/l									
Chromium VI	EPA 7196A	7C08171	0.65	25	4.7	1	03/08/07	03/08/07	J
Sample ID: IQC0980-10 (MW3010_WG030807_0001 - Water)					Sampled: 03/08/07				
Reporting Units: ug/l									
Chromium VI	EPA 7196A	7C08171	3.2	120	140	5	03/08/07	03/08/07	

TestAmerica - Irvine, CA
Nicholas Marz
Project Manager

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The results pertain only to the samples tested in the laboratory. This report shall not be reproduced, except in full, without written permission from TestAmerica.

IQC0980 <Page 2 of 7>

LDC #: 16591A6
 SDG #: IQC0980
 Laboratory: Test America

VALIDATION COMPLETENESS WORKSHEET
 Tier 1/2/3

Date: 4/21/07
 Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: Hexavalent Chromium (EPA SW846 Method 7196A)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: <u>3/8/07</u>
IIa.	Initial calibration	A	Not reviewed for Tier I validation.
IIb.	Calibration verification	A	Not reviewed for Tier I validation.
III.	Blanks	A	
IVa.	Matrix Spike/(Matrix Spike) Duplicates	A	<u>MS/MSD IQC0980-0</u>
IVb.	Laboratory control samples	A	<u>LC5</u>
V.	Sample result verification	A	Not reviewed for Tier I or Tier II validation.
VI.	Overall assessment of data	A	
VII.	Field duplicates	<u>SW</u>	<u>(2,3)</u>
VIII.	Field blanks	<u>N</u>	

Note: A = Acceptable ND = No compounds detected D = Duplicate
 N = Not provided/applicable R = Rinsate TB = Trip blank
 SW = See worksheet FB = Field blank EB = Equipment blank

Validated Samples: * Indicates sample underwent Tier I validation, ** Indicates sample underwent Tier III validation

1	MW3009_WG030807_0001 ^A	11		21		31	
2	MW3012_WG030807_0001 [*]	12		22		32	
3	MW3012_WG030807_0002 ^{**}	13		23		33	
4	<u>VB</u>	14		24		34	
5		15		25		35	
6		16		26		36	
7		17		27		37	
8		18		28		38	
9		19		29		39	
10		20		30		40	

Notes: _____

LDC #: 16591A6
 SDG #: See com

VALIDATION FINDINGS CHECKLIST

Page: 1 of 1
 Reviewer: MY
 2nd Reviewer: [Signature]

Method: Inorganics (EPA Method 7196A)

Validation Area	Yes	No	NA	Findings/Comments
All technical holding times were met.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Cooler temperature criteria was met.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all instruments calibrated daily, each set-up time?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the proper number of standards used?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all initial calibration correlation coefficients > 0.995?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all initial and continuing calibration verification %Rs within the 90-110% QC limits?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were titrant checks performed as required? (Level IV only)	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were balance checks performed as required? (Level IV only)	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Was a method blank associated with every sample in this SDG?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was there contamination in the method blanks? If yes, please see the Blanks validation completeness worksheet.	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
Were a matrix spike (MS) and duplicate (DUP) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD or MS/DUP. Soil / Water.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the 75-125 QC limits? If the sample concentration exceeded the spike concentration by a factor of 4 or more, no action was taken.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the MS/MSD or duplicate relative percent differences (RPD) ≤ 20% for waters and ≤ 35% for soil samples? A control limit of ≤ CRDL (≤ 2X CRDL for soil) was used for samples that were ≤ 5X the CRDL, including when only one of the duplicate sample values were < 5X the CRDL.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was an LCS analyzed for this SDG?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was an LCS analyzed per extraction batch?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the 80-120% (85-115% for Method 300.0) QC limits?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were performance evaluation (PE) samples performed?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were the performance evaluation (PE) samples within the acceptance limits?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	

LDC #: 16591166
 SDG #: see con

VALIDATION FINDINGS CHECKLIST

Page: 1 of 1
 Reviewer: WY
 2nd Reviewer: [Signature]

Validation Area	Yes	No	NA	Findings/Comments
Were RLs adjusted to reflect all sample dilutions and dry weight factors applicable to level IV validation?	✓			
Were detection limits < RL?	✓			
Overall assessment of data was found to be acceptable.	✓			
Field duplicate pairs were identified in this SDG.	✓			
Target analytes were detected in the field duplicates.	✓			
Field blanks were identified in this SDG.			✓	
Target analytes were detected in the field blanks.			✓	

LDC#: 16591A6
SDG#: IQC0980

VALIDATION FINDINGS WORKSHEET
Field Duplicates

Page: (of)
Reviewer: [Signature]
2nd Reviewer: [Signature]

Inorganics, Method 7196A

- Y N NA Were field duplicate pairs identified in this SDG?
- Y N NA Were target analytes detected in the field duplicate pairs?

Analyte	Concentration (ug/L)		RPD	
	2	3		
Cr (VI)	1.3	0.98	28	

V:\FIELD DUPLICATES\FD_inorganic16591A6.wpd

LDC #: 16811A6
 SDG #: See cover

VALIDATION FINDINGS WORKSHEET
Initial and Continuing Calibration Calculation Verification

Page: 1 of 1
 Reviewer: HW
 2nd Reviewer: AL

METHOD: Inorganics, Method 7196A
 The correlation coefficient (r) for the calibration of Cu⁶⁺ was recalculated. Calibration date: 3/8/07

An initial or continuing calibration verification percent recovery (%R) was recalculated for each type of analysis using the following formula:

$\%R = \frac{\text{Found}}{\text{True}} \times 100$ Where, Found = concentration of each analyte measured in the analysis of the ICV or CCV solution
 True = concentration of each analyte in the ICV or CCV source

Type of Analysis	Analyte	Concn. (ug/L) (units)	Concn. (ug/L) (units)	Recalculated		Acceptable (Y/N)	
				r or %R	r or %R		
Initial calibration		Blank	0				
Calibration verification	Cu ⁶⁺	Standard 1	0.07	0.007			
		Standard 2	0.025	0.023			
		Standard 3	0.1	0.083			
		Standard 4	0.5	0.414	$r = 0.99997$	$r = 0.99998$	Y
		Standard 5					
		Standard 6					
		Standard 7					
Calibration verification ICV	Cu ⁶⁺	0.1	0.101	101	NR	Y	
Calibration verification CCV	Cu ⁶⁺	0.3	0.310	103	NR	Y	
Calibration verification							

Comments: Refer to Calibration Verification findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 16091A6
 SDG #: Lee

VALIDATION FINDINGS WORKSHEET
Level IV Recalculation Worksheet

Page: 1 of 1
 Reviewer: MM
 2nd Reviewer: R

METHOD: Inorganics, Method 9196A

Percent recoveries (%R) for a laboratory control sample and a matrix spike sample were recalculated using the following formula:

$$\%R = \frac{\text{Found}}{\text{True}} \times 100$$
 Where, Found = concentration of each analyte measured in the analysis of the sample. For the matrix spike calculation, Found = SSR (spiked sample result) - SR (sample result).
 True = concentration of each analyte in the source.

A sample and duplicate relative percent difference (RPD) was recalculated using the following formula:

$$RPD = \frac{|S-D|}{(S+D)/2} \times 100$$
 Where, S = Original sample concentration
 D = Duplicate sample concentration

Sample ID	Type of Analysis	Element	Found / S (units)	True / D (units)	Recalculated		Reported		Acceptable (Y/N)
					%R / RPD	%R / RPD			
<u>LS</u>	Laboratory control sample	<u>Cr</u>	<u>101</u>	<u>100</u>	<u>101</u>	<u>101</u>			<u>Y</u>
<u>IPAC-980-101</u>	Matrix spike sample	<u>✓</u>	<u>305</u> (SSR-SR)	<u>300</u>	<u>102</u>	<u>102</u>			<u>Y</u>
<u>✓</u>	Duplicate sample		<u>304</u>	<u>305</u>	<u>0</u>	<u>0</u>			<u>Y</u>

Comments: Refer to appropriate worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

**Boeing Realty Corp., Bldg C-1 Long Beach
Data Validation Reports
LDC# 16591**

TPH as Extractables

LDC

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Boeing Realty Corp., Bldg C-1 Long Beach
Collection Date: March 15, 2007
LDC Report Date: April 30, 2007
Matrix: Water
Parameters: Total Petroleum Hydrocarbons as Extractables
Validation Level: Tier 3
Laboratory: TestAmerica
Sample Delivery Group (SDG): IQC1776
Sample Identification
MW3017_WG031507_0001

Introduction

This data review covers one water sample listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 8015 for Total Petroleum Hydrocarbons (TPH) as Extractables.

This review follows a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Organic Data Review (October 1999) as there are no current guidelines for the method stated above.

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

Blank results are summarized in Section III.

Field duplicates are summarized in Section IX.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- N Presumptive evidence of presence of the constituent.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. Calibration

a. Initial Calibration

Initial calibration of compounds was performed as required by the method.

The percent relative standard deviations (%RSD) of calibration factors for compounds were less than or equal to 20.0% .

b. Calibration Verification

Calibration verification was performed at required frequencies. The percent differences (%D) of amounts in continuing standard mixtures were within the 15.0% QC limits.

The percent difference (%D) of the second source calibration standard were less than or equal to 25.0% for all compounds.

III. Blanks

Method blanks were reviewed for each matrix as applicable. No total petroleum hydrocarbons as extractable contaminants were found in the method blanks.

IV. Accuracy and Precision Data

a. Surrogate Recovery

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

b. Matrix Spike/Matrix Spike Duplicates

The laboratory has indicated that there were no matrix spike (MS) and matrix spike duplicate (MSD) analyses specified for the samples in this SDG, and therefore matrix spike and matrix spike duplicate analyses were not performed for this SDG.

c. Laboratory Control Samples

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

V. Target Compound Identification

All target compound identifications were within validation criteria.

VI. Compound Quantitation and CRQLs

All compound quantitation and CRQLs were within validation criteria.

VII. System Performance

The system performance was acceptable.

VIII. Overall Assessment of Data

Data flags have been summarized at the end of this report if data has been summarized.

IX. Field Duplicates

No field duplicates were identified in this SDG.

X. Field Blanks

No field blanks were identified in this SDG.

**Boeing Realty Corp., Bldg C-1 Long Beach
Total Petroleum Hydrocarbons as Extractables - Data Qualification Summary - SDG
IQC1776**

No Sample Data Qualified in this SDG

**Boeing Realty Corp., Bldg C-1 Long Beach
Total Petroleum Hydrocarbons as Extractables - Laboratory Blank Data Qualification
Summary - SDG IQC1776**

No Sample Data Qualified in this SDG

TestAmerica

ANALYTICAL TESTING CORPORATION

17461 Derian Avenue, Suite 100, Irvine, CA 92614 (949) 261-1022 Fax: (949) 260-3297

TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-1 Long Beach
EM-2701
Report Number: IQC1776

Sampled: 03/15/07
Received: 03/15/07

HYDROCARBON DISTRIBUTION (EPA 3510C/8015 Mod.)

Analyte	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	% of Total	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC1776-04 (MW3017_WG031507_0001 - Water)									
Reporting Units: mg/l									
EFH (C6 - C44)	7C22062	0.094	0.47	4.9	1	100	3/22/2007	3/22/2007	
EFH (C6 - C7)	7C22062	0.094	0.094	ND	1	N/A	3/22/2007	3/22/2007	
EFH (C8 - C9)	7C22062	0.094	0.094	0.15	1	3	3/22/2007	3/22/2007	
EFH (C10 - C11)	7C22062	0.094	0.094	0.79	1	16	3/22/2007	3/22/2007	
EFH (C12 - C13)	7C22062	0.094	0.094	1.1	1	22	3/22/2007	3/22/2007	
EFH (C14 - C15)	7C22062	0.094	0.094	1.4	1	29	3/22/2007	3/22/2007	
EFH (C16 - C17)	7C22062	0.094	0.094	0.93	1	19	3/22/2007	3/22/2007	
EFH (C18 - C19)	7C22062	0.094	0.094	0.29	1	6	3/22/2007	3/22/2007	
EFH (C20 - C23)	7C22062	0.042	0.042	0.084	1	2	3/22/2007	3/22/2007	
EFH (C24 - C27)	7C22062	0.042	0.042	0.045	1	1	3/22/2007	3/22/2007	
EFH (C28 - C31)	7C22062	0.042	0.042	ND	1	N/A	3/22/2007	3/22/2007	
EFH (C32 - C35)	7C22062	0.094	0.094	ND	1	N/A	3/22/2007	3/22/2007	
EFH (C36 - C39)	7C22062	0.042	0.042	ND	1	N/A	3/22/2007	3/22/2007	
EFH (C40 - C44)	7C22062	0.042	0.042	ND	1	N/A	3/22/2007	3/22/2007	
Surrogate: n-Octacosane (40-125%)				97%					

TestAmerica - Irvine, CA
Nicholas Marz
Project Manager

Handwritten signature

The results pertain only to the samples tested in the laboratory. This report shall not be reproduced, except in full, without written permission from TestAmerica.

IQC1776 <Page 18 of 37>

LDC #: 16591B8
 SDG #: IQC1776
 Laboratory: Test America

VALIDATION COMPLETENESS WORKSHEET
 EPA Region 1 - Tier 3

Date: 4/25/07
 Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: GC TPH as Extractables (EPA SW 846 Method 8015)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: 3/15/07
IIa.	Initial calibration	A	
IIb.	Calibration verification	A	
III.	Blanks	A	
IVa.	Surrogate recovery	A	
IVb.	Matrix spike/Matrix spike duplicates	N	chem specified
IVc.	Laboratory control samples	A	res ID
V.	Target compound identification	A	
VI.	Compound Quantitation and CRQLs	A	
VII.	System Performance	A	
VIII.	Overall assessment of data	A	
IX.	Field duplicates	N	
X.	Field blanks	N	

Note: A = Acceptable ND = No compounds detected D = Duplicate
 N = Not provided/applicable R = Rinsate TB = Trip blank
 SW = See worksheet FB = Field blank EB = Equipment blank

Validated Samples: *wait*

1	MW3017-WG031507_0001	11	TC22062-BLK1	21		31	
2		12		22		32	
3		13		23		33	
4		14		24		34	
5		15		25		35	
6		16		26		36	
7		17		27		37	
8		18		28		38	
9		19		29		39	
10		20		30		40	

Notes: _____

LDC #: 1659138
 SDG #: per cover

VALIDATION FINDINGS CHECKLIST

Page: 1 of 2
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

Method: GC HPLC

Validation Area	Yes	No	NA	Findings/Comments
All technical holding times were met.	/			
Cooler temperature criteria was met.	/			
Did the laboratory perform a 5 point calibration prior to sample analysis?	/			
Was a linear fit used for evaluation? If yes, were all percent relative standard deviations (%RSD) < 20%?	/			
Was a curve fit used for evaluation? If Yes, what was the acceptance criteria used?			/	
Did the initial calibration meet the curve fit acceptance criteria?			/	
Were the RT windows properly established?	/			
What type of continuing calibration calculation was performed? ___%D or %R	/			
Was a continuing calibration analyzed daily?	/			
Were all percent differences (%D) < 15%.0 or percent recoveries 85-115%?	/			
Were all the retention times within the acceptance windows?	/			
Was a method blank associated with every sample in this SDG?	/			
Was a method blank analyzed for each matrix and concentration?			/	
Was there contamination in the method blanks? If yes, please see the Blanks validation completeness worksheet.			/	
Were all surrogate %R within the QC limits?	/			
If the percent recovery (%R) of one or more surrogates was outside QC limits, was a reanalysis performed to confirm %R?			/	
If any %R was less than 10 percent, was a reanalysis performed to confirm %R?			/	
Were a matrix spike (MS) and matrix spike duplicate (MSD) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD. Soil / Water.			/	
Was a MS/MSD analyzed every 20 samples of each matrix?			/	
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the QC limits?			/	
Was an LCS analyzed for this SDG?	/			
Was an LCS analyzed per extraction batch?	/			

LDC #: 16591B8
 SDG #: all cover

VALIDATION FINDINGS CHECKLIST

Page: 2 of 4
 Reviewer: 15
 2nd Reviewer: A

Validation Area	Yes	No	NA	Findings/Comments
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the QC limits?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were performance evaluation (PE) samples performed?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were the performance evaluation (PE) samples within the acceptance limits?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were the retention times of reported detects within the RT windows?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were compound quantitation and CRQLs adjusted to reflect all sample dilutions and dry weight factors applicable to level IV validation?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
System performance was found to be acceptable.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Overall assessment of data was found to be acceptable.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were field duplicate pairs identified in this SDG?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were target compounds idetected in the field duplicates?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were field blanks identified in this SDG?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were target compounds detected in the field blanks?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	

LDC #: 1659188
 SDG #: see cover

VALIDATION FINDINGS WORKSHEET
Initial Calibration Calculation Verification

Page: 1 of 1
 Reviewer: R
 2nd Reviewer:

METHOD: GC HPLC

The calibration Factor (CF), average CF, and percent relative standard deviation (%RSD) were recalculated for the compounds identified below using the following calculations:

- CF = A/C
- average CF = sum of the CF/number of standards
- %RSD = $100 \cdot (S/X)$
- A = Area of compound,
- C = Concentration of compound,
- S = Standard deviation of the CF
- X = Mean of the CFs

#	Standard ID	Calibration Date	Compound	Reported		Recalculated		Reported		Recalculated	
				CF (1SD std)	CF (1SD std)	Average CF (Initial)	Average CF (Initial)	%RSD	%RSD		
1	1CAL	3/3/07	EFH	2873.4	2473.4	2787.17	2787.17	11.14	11.14		
2											
3											
4											

Comments: Refer to Initial Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

VALIDATION FINDINGS WORKSHEET
Surrogate Results Verification

The percent recoveries (%R) of surrogates were recalculated for the compounds identified below using the following calculation:

% Recovery: $SF/SS \cdot 100$ Where: SF = Surrogate Found
 SS = Surrogate Spiked

Sample ID: #

Surrogate	Column/Detector	Surrogate Spiked	Surrogate Found	Percent Recovery		Percent Difference
				Reported	Recalculated	
<u>n-octadecane</u>	<u>not spiked</u>	<u>100</u>	<u>97.4363</u>	<u>97</u>	<u>97</u>	<u>0</u>

Sample ID: _____

Surrogate	Column/Detector	Surrogate Spiked	Surrogate Found	Percent Recovery		Percent Difference
				Reported	Recalculated	

Sample ID: _____

Surrogate	Column/Detector	Surrogate Spiked	Surrogate Found	Percent Recovery		Percent Difference
				Reported	Recalculated	

LDC #: 1659138 VALIDATION FINDINGS WORKSHEET

Page: 61

SDG #: per cover Laboratory Control Sample/Laboratory Control Sample Duplicates Results Verification

Reviewer: [Signature]

2nd Reviewer: [Signature]

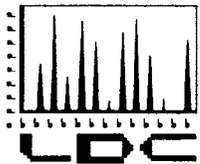
METHOD: GC HPLC

The percent recoveries (%R) and relative percent differences (RPD) of the laboratory control sample and laboratory control sample duplicate were recalculated for the compounds identified below using the following calculation:

%Recovery = $100 \cdot (SSC - SC) / SA$ Where SSC = Spiked sample concentration SC = Sample concentration
 SA = Spike added
 RPD = $(|(SSCLCS - SSCLCSD) \cdot 2| / (SSCLCS + SSCLCSD)) \cdot 100$ LCS = Laboratory Control Sample LCSD = Laboratory Control Sample duplicate
 LCS/LCSD samples: 1022062-135

Compound	Spike Added (mg/L)		Sample Conc. (mg/L)	Spike Sample Concentration (mg/L)		LCS		LCSD		LCS/LCSD	
	LCS	LCSD		LCS	LCSD	Percent Recovery	Recalc.	Percent Recovery	Recalc.	Reported	Recalc.
Gasoline (8015)											
Diesel (8015)											
Benzene (8021B)											
Methane (RSK-175)											
2,4-D (8151)											
Dinoseb (8151)											
Naphthalene (8310)											
Anthracene (8310)											
HMX (8330)											
2,4,6-Trinitrotoluene (8330)											
EFH	1.0	1.0	0	0.931	0.806	93	93	81	81	14	14

Comments: Refer to Laboratory Control Sample/Laboratory Control Sample Duplicate findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.



LABORATORY DATA CONSULTANTS, INC.

7750 El Camino Real, Suite 2L Carlsbad, CA 92009 Phone: 760/634-0437 Fax: 760/634-0439

Tait Environmental Management, Inc.
701 N. Park Center Drive
Santa Ana, CA 92705
ATTN: Ms. Clara Boeru

May 11, 2007

SUBJECT: Boeing Realty Corp. Bldg C-6 Torrance, Data Validation

Dear Ms. Boeru,

Enclosed are the final validation reports for the fractions listed below. These SDGs were received on May 3, 2007. Attachment 1 is a summary of the samples that were reviewed for each analysis.

LDC Project # 16739:

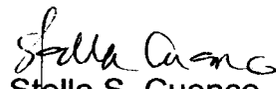
<u>SDG #</u>	<u>Fraction</u>
IQC1612, IQC2470, IQC2895	Volatiles, Manganese, Wet Chemistry, Dissolved Gases

The data validation was performed under Tier 1, Tier 2 and Tier 3 guidelines. The analyses were validated using the following documents, as applicable to each method:

- USEPA, Contract Laboratory Program National Functional Guidelines for Organic Data Review, October 1999
- USEPA, Contract Laboratory Program National Functional Guidelines for Inorganic Data Review, October 1999
- EPA SW 846, Third Edition, Test Methods for Evaluating Solid Waste, update 1, July 1992; update IIA, August 1993; update II, September 1994; update IIB, January 1995; update III, December 1996; update IIIA, April 1998

Please feel free to contact us if you have any questions.

Sincerely,


Stella S. Cuenco
Project Manager/Senior Chemist

LDC #16739 (Tait Environmental Management, Inc. / Boeing Realty Corp., Bldg C-6 Torrance)

LDC	SDG#	DATE REC'D	(3) DATE DUE	VOA (8260B)		Mh (6010B)		Diss. Gases (175)		Alk. (310.1)		NH ₃ (350.3)		Cl,SO ₄ , O-PO ₄ (300.0)		NO ₂ -N, NO ₃ -N (300.0)		S= (376.2)		TOC (415.1)		Tier I		Tier I					
				W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S
A	IQC1612	05/03/07	05/24/07	6	0	2	0	2	0	2	0	2	0	2	0	2	0	2	0	-	-	2	0						
B	IQC2470	05/03/07	05/24/07	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	1	0	0	0							
B	IQC2470	05/03/07	05/24/07	4	0	1	0	1	0	1	0	1	0	1	0	1	0	1	0	1	0	1							
C	IQC2895	05/03/07	05/24/07	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0							
				11	0	4	0	4	0	4	0	4	0	4	0	4	0	4	0	3	0	4	0	0	0	0	0	0	0
Total																													38

Shaded cells indicate Tier III validation (all other cells are Tier II validation). Sample counts do not include MS, MSD, or DUP's.

**Boeing Realty Corp., Bldg C-6 Torrance
Data Validation Reports
LDC# 16739**

Volatiles

LDC

LDC Report# 16739A1

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Boeing Realty Corp., Bldg C-6 Torrance
Collection Date: March 14, 2007
LDC Report Date: May 10, 2007
Matrix: Water
Parameters: Volatiles
Validation Level: Tier 1
Laboratory: TestAmerica

Sample Delivery Group (SDG): IQC1612

Sample Identification

MWB013_WG031407_0001
MWG004_WG031407_0001
TMW_14_WG031407_0001
TMW_11_WG031407_0001
WCC_5S_WG031407_0001
MWC021_WG031407_0001
MWB013_WG031407_0001MS
MWB013_WG031407_0001MSD

Introduction

This data review covers 8 water samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 8260B for Volatiles.

This review follows a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Organic Data Review (October 1999) as there are no current guidelines for the method stated above.

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

Blank results are summarized in Section V.

Field duplicates are summarized in Section XVI.

Raw data were not reviewed for this SDG. The review was based on QC data.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- N Presumptive evidence of presence of the constituent.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. GC/MS Instrument Performance Check

Instrument performance data were not reviewed for Tier 1.

III. Initial Calibration

Initial calibration data were not reviewed for Tier 1.

IV. Continuing Calibration

Continuing calibration data were not reviewed for Tier 1.

V. Blanks

Method blanks were reviewed for each matrix as applicable. No volatile contaminants were found in the method blanks.

VI. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VII. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

VIII. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) were within QC limits.

IX. Regional Quality Assurance and Quality Control

Not applicable.

X. Internal Standards

Internal standards data were not reviewed for Tier 1.

XI. Target Compound Identifications

Raw data were not reviewed for this SDG.

XII. Compound Quantitation and CRQLs

Raw data were not reviewed for this SDG.

XIII. Tentatively Identified Compounds (TICs)

Raw data were not reviewed for this SDG.

XIV. System Performance

Raw data were not reviewed for this SDG.

XV. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

XVI. Field Duplicates

No field duplicates were identified in this SDG.

XVII. Field Blanks

No field blanks were identified in this SDG.

**Boeing Realty Corp., Bldg C-6 Torrance
Volatiles - Data Qualification Summary - SDG IQC1612**

No Sample Data Qualified in this SDG

**Boeing Realty Corp., Bldg C-6 Torrance
Volatiles - Laboratory Blank Data Qualification Summary - SDG IQC1612**

No Sample Data Qualified in this SDG

TestAmerica

ANALYTICAL TESTING CORPORATION

17461 Derian Avenue, Suite 100, Irvine, CA 92614 (949) 261-1022 Fax: (949) 260-3297

TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727-01
Report Number: IQC1612

Sampled: 03/14/07
Received: 03/14/07

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC1612-05 (MWB013_WG031407_0001 - Water)									
Reporting Units: ug/l									
Acetone	EPA 8260B	7C16022	4.5	10	ND	1	03/16/07	03/16/07	
Benzene	EPA 8260B	7C16022	0.28	1.0	ND	1	03/16/07	03/16/07	
Bromobenzene	EPA 8260B	7C16022	0.27	1.0	ND	1	03/16/07	03/16/07	
Bromochloromethane	EPA 8260B	7C16022	0.32	1.0	ND	1	03/16/07	03/16/07	
Bromodichloromethane	EPA 8260B	7C16022	0.30	1.0	0.37	1	03/16/07	03/16/07	J
Bromoform	EPA 8260B	7C16022	0.40	1.0	ND	1	03/16/07	03/16/07	
Bromomethane	EPA 8260B	7C16022	0.42	1.0	ND	1	03/16/07	03/16/07	
2-Butanone (MEK)	EPA 8260B	7C16022	3.8	5.0	ND	1	03/16/07	03/16/07	
n-Butylbenzene	EPA 8260B	7C16022	0.37	1.0	ND	1	03/16/07	03/16/07	
sec-Butylbenzene	EPA 8260B	7C16022	0.25	1.0	ND	1	03/16/07	03/16/07	
tert-Butylbenzene	EPA 8260B	7C16022	0.22	1.0	ND	1	03/16/07	03/16/07	
Carbon Disulfide	EPA 8260B	7C16022	0.48	1.0	ND	1	03/16/07	03/16/07	
Carbon tetrachloride	EPA 8260B	7C16022	0.28	0.50	ND	1	03/16/07	03/16/07	
Chlorobenzene	EPA 8260B	7C16022	0.36	1.0	ND	1	03/16/07	03/16/07	
Chloroethane	EPA 8260B	7C16022	0.40	2.0	ND	1	03/16/07	03/16/07	
Chloroform	EPA 8260B	7C16022	0.33	1.0	0.54	1	03/16/07	03/16/07	J
Chloromethane	EPA 8260B	7C16022	0.40	2.0	ND	1	03/16/07	03/16/07	
2-Chlorotoluene	EPA 8260B	7C16022	0.28	1.0	ND	1	03/16/07	03/16/07	
4-Chlorotoluene	EPA 8260B	7C16022	0.29	1.0	ND	1	03/16/07	03/16/07	
1,2-Dibromo-3-chloropropane	EPA 8260B	7C16022	0.97	2.0	ND	1	03/16/07	03/16/07	
Dibromochloromethane	EPA 8260B	7C16022	0.28	1.0	ND	1	03/16/07	03/16/07	
1,2-Dibromoethane (EDB)	EPA 8260B	7C16022	0.40	1.0	ND	1	03/16/07	03/16/07	
1,4-Dichlorobenzene	EPA 8260B	7C16022	0.37	1.0	ND	1	03/16/07	03/16/07	
1,2-Dichlorobenzene	EPA 8260B	7C16022	0.32	1.0	ND	1	03/16/07	03/16/07	
1,3-Dichlorobenzene	EPA 8260B	7C16022	0.35	1.0	ND	1	03/16/07	03/16/07	
Dichlorodifluoromethane	EPA 8260B	7C16022	0.79	1.0	ND	1	03/16/07	03/16/07	
1,2-Dichloroethane	EPA 8260B	7C16022	0.28	0.50	ND	1	03/16/07	03/16/07	
1,1-Dichloroethane	EPA 8260B	7C16022	0.27	1.0	ND	1	03/16/07	03/16/07	
1,1-Dichloroethene	EPA 8260B	7C16022	0.42	1.0	ND	1	03/16/07	03/16/07	
cis-1,2-Dichloroethene	EPA 8260B	7C16022	0.32	1.0	ND	1	03/16/07	03/16/07	
trans-1,2-Dichloroethene	EPA 8260B	7C16022	0.27	1.0	ND	1	03/16/07	03/16/07	
1,2-Dichloropropane	EPA 8260B	7C16022	0.35	1.0	ND	1	03/16/07	03/16/07	
2,2-Dichloropropane	EPA 8260B	7C16022	0.34	1.0	ND	1	03/16/07	03/16/07	
cis-1,3-Dichloropropene	EPA 8260B	7C16022	0.22	0.50	ND	1	03/16/07	03/16/07	
1,1-Dichloropropene	EPA 8260B	7C16022	0.28	1.0	ND	1	03/16/07	03/16/07	
trans-1,3-Dichloropropene	EPA 8260B	7C16022	0.32	0.50	ND	1	03/16/07	03/16/07	
Ethylbenzene	EPA 8260B	7C16022	0.25	1.0	ND	1	03/16/07	03/16/07	
Hexachlorobutadiene	EPA 8260B	7C16022	0.38	1.0	ND	1	03/16/07	03/16/07	
2-Hexanone	EPA 8260B	7C16022	2.6	6.0	ND	1	03/16/07	03/16/07	
Iodomethane	EPA 8260B	7C16022	1.0	2.0	ND	1	03/16/07	03/16/07	
Isopropylbenzene	EPA 8260B	7C16022	0.25	1.0	ND	1	03/16/07	03/16/07	

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Nicholas Marz
Project Manager

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TestAmerica

ANALYTICAL TESTING CORPORATION

17461 Derian Avenue, Suite 100, Irvine, CA 92614 (949) 261-1022 Fax: (949) 260-3297

TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727-01
Report Number: IQC1612

Sampled: 03/14/07
Received: 03/14/07

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC1612-05 (MWB013_WG031407_0001 - Water) - cont.									
Reporting Units: ug/l									
p-Isopropyltoluene	EPA 8260B	7C16022	0.28	1.0	ND	1	03/16/07	03/16/07	
Methyl-tert-butyl Ether (MTBE)	EPA 8260B	7C16022	0.32	1.0	ND	1	03/16/07	03/16/07	
Methylene chloride	EPA 8260B	7C16022	0.95	1.0	ND	1	03/16/07	03/16/07	
4-Methyl-2-pentanone (MIBK)	EPA 8260B	7C16022	3.5	5.0	ND	1	03/16/07	03/16/07	
n-Propylbenzene	EPA 8260B	7C16022	0.27	1.0	ND	1	03/16/07	03/16/07	
Styrene	EPA 8260B	7C16022	0.16	1.0	ND	1	03/16/07	03/16/07	
1,1,1,2-Tetrachloroethane	EPA 8260B	7C16022	0.27	1.0	ND	1	03/16/07	03/16/07	
1,1,2,2-Tetrachloroethane	EPA 8260B	7C16022	0.24	1.0	ND	1	03/16/07	03/16/07	
Tetrachloroethene	EPA 8260B	7C16022	0.32	1.0	ND	1	03/16/07	03/16/07	
Tetrahydrofuran (THF)	EPA 8260B	7C16022	3.5	10	ND	1	03/16/07	03/16/07	
Toluene	EPA 8260B	7C16022	0.36	1.0	ND	1	03/16/07	03/16/07	
1,2,3-Trichlorobenzene	EPA 8260B	7C16022	0.30	1.0	ND	1	03/16/07	03/16/07	
1,2,4-Trichlorobenzene	EPA 8260B	7C16022	0.48	1.0	ND	1	03/16/07	03/16/07	
1,1,2-Trichloroethane	EPA 8260B	7C16022	0.30	1.0	ND	1	03/16/07	03/16/07	
1,1,1-Trichloroethane	EPA 8260B	7C16022	0.30	1.0	ND	1	03/16/07	03/16/07	
Trichloroethene	EPA 8260B	7C16022	0.26	1.0	5.3	1	03/16/07	03/16/07	
Trichlorofluoromethane	EPA 8260B	7C16022	0.34	2.0	ND	1	03/16/07	03/16/07	
1,2,3-Trichloropropane	EPA 8260B	7C16022	0.40	1.0	ND	1	03/16/07	03/16/07	
1,2,4-Trimethylbenzene	EPA 8260B	7C16022	0.23	1.0	ND	1	03/16/07	03/16/07	
1,3,5-Trimethylbenzene	EPA 8260B	7C16022	0.26	1.0	ND	1	03/16/07	03/16/07	
Vinyl acetate	EPA 8260B	7C16022	1.7	6.0	ND	1	03/16/07	03/16/07	
Vinyl chloride	EPA 8260B	7C16022	0.30	0.50	ND	1	03/16/07	03/16/07	
Xylenes, Total	EPA 8260B	7C16022	0.90	1.0	ND	1	03/16/07	03/16/07	
Surrogate: 4-Bromofluorobenzene (80-120%)					108 %				
Surrogate: Dibromofluoromethane (80-120%)					109 %				
Surrogate: Toluene-d8 (80-120%)					103 %				

TestAmerica - Irvine, CA
Nicholas Marz
Project Manager

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ANALYTICAL TESTING CORPORATION

17461 Derian Avenue, Suite 100, Irvine, CA 92614 (949) 261-1022 Fax: (949) 260-3297

TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727-01
Report Number: IQC1612

Sampled: 03/14/07
Received: 03/14/07

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC1612-06 (MWG004_WG031407_0001 - Water)									
Reporting Units: ug/l									
Acetone	EPA 8260B	7C16022	4.5	10	ND	1	03/16/07	03/16/07	
Benzene	EPA 8260B	7C16022	0.28	1.0	ND	1	03/16/07	03/16/07	
Bromobenzene	EPA 8260B	7C16022	0.27	1.0	ND	1	03/16/07	03/16/07	
Bromochloromethane	EPA 8260B	7C16022	0.32	1.0	ND	1	03/16/07	03/16/07	
Bromodichloromethane	EPA 8260B	7C16022	0.30	1.0	ND	1	03/16/07	03/16/07	
Bromoform	EPA 8260B	7C16022	0.40	1.0	ND	1	03/16/07	03/16/07	
Bromomethane	EPA 8260B	7C16022	0.42	1.0	ND	1	03/16/07	03/16/07	
2-Butanone (MEK)	EPA 8260B	7C16022	3.8	5.0	ND	1	03/16/07	03/16/07	
n-Butylbenzene	EPA 8260B	7C16022	0.37	1.0	ND	1	03/16/07	03/16/07	
sec-Butylbenzene	EPA 8260B	7C16022	0.25	1.0	ND	1	03/16/07	03/16/07	
tert-Butylbenzene	EPA 8260B	7C16022	0.22	1.0	ND	1	03/16/07	03/16/07	
Carbon Disulfide	EPA 8260B	7C16022	0.48	1.0	ND	1	03/16/07	03/16/07	
Carbon tetrachloride	EPA 8260B	7C16022	0.28	0.50	ND	1	03/16/07	03/16/07	
Chlorobenzene	EPA 8260B	7C16022	0.36	1.0	ND	1	03/16/07	03/16/07	
Chloroethane	EPA 8260B	7C16022	0.40	2.0	ND	1	03/16/07	03/16/07	
Chloroform	EPA 8260B	7C16022	0.33	1.0	ND	1	03/16/07	03/16/07	
Chloromethane	EPA 8260B	7C16022	0.40	2.0	ND	1	03/16/07	03/16/07	
2-Chlorotoluene	EPA 8260B	7C16022	0.28	1.0	ND	1	03/16/07	03/16/07	
4-Chlorotoluene	EPA 8260B	7C16022	0.29	1.0	ND	1	03/16/07	03/16/07	
1,2-Dibromo-3-chloropropane	EPA 8260B	7C16022	0.97	2.0	ND	1	03/16/07	03/16/07	
Dibromochloromethane	EPA 8260B	7C16022	0.28	1.0	ND	1	03/16/07	03/16/07	
1,2-Dibromoethane (EDB)	EPA 8260B	7C16022	0.40	1.0	ND	1	03/16/07	03/16/07	
1,4-Dichlorobenzene	EPA 8260B	7C16022	0.37	1.0	ND	1	03/16/07	03/16/07	
1,2-Dichlorobenzene	EPA 8260B	7C16022	0.32	1.0	ND	1	03/16/07	03/16/07	
1,3-Dichlorobenzene	EPA 8260B	7C16022	0.35	1.0	ND	1	03/16/07	03/16/07	
Dichlorodifluoromethane	EPA 8260B	7C16022	0.79	1.0	ND	1	03/16/07	03/16/07	
1,2-Dichloroethane	EPA 8260B	7C16022	0.28	0.50	ND	1	03/16/07	03/16/07	
1,1-Dichloroethane	EPA 8260B	7C16022	0.27	1.0	ND	1	03/16/07	03/16/07	
1,1-Dichloroethene	EPA 8260B	7C16022	0.42	1.0	ND	1	03/16/07	03/16/07	
cis-1,2-Dichloroethene	EPA 8260B	7C16022	0.32	1.0	0.34	1	03/16/07	03/16/07	J
trans-1,2-Dichloroethene	EPA 8260B	7C16022	0.27	1.0	ND	1	03/16/07	03/16/07	
1,2-Dichloropropane	EPA 8260B	7C16022	0.35	1.0	ND	1	03/16/07	03/16/07	
2,2-Dichloropropane	EPA 8260B	7C16022	0.34	1.0	ND	1	03/16/07	03/16/07	
cis-1,3-Dichloropropene	EPA 8260B	7C16022	0.22	0.50	ND	1	03/16/07	03/16/07	
1,1-Dichloropropene	EPA 8260B	7C16022	0.28	1.0	ND	1	03/16/07	03/16/07	
trans-1,3-Dichloropropene	EPA 8260B	7C16022	0.32	0.50	ND	1	03/16/07	03/16/07	
Ethylbenzene	EPA 8260B	7C16022	0.25	1.0	ND	1	03/16/07	03/16/07	
Hexachlorobutadiene	EPA 8260B	7C16022	0.38	1.0	ND	1	03/16/07	03/16/07	
2-Hexanone	EPA 8260B	7C16022	2.6	6.0	ND	1	03/16/07	03/16/07	
Iodomethane	EPA 8260B	7C16022	1.0	2.0	ND	1	03/16/07	03/16/07	
Isopropylbenzene	EPA 8260B	7C16022	0.25	1.0	ND	1	03/16/07	03/16/07	

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Nicholas Marz
Project Manager

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ANALYTICAL TESTING CORPORATION

17461 Derian Avenue, Suite 100, Irvine, CA 92614 (949) 261-1022 Fax: (949) 260-3297

TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727-01
Report Number: IQC1612

Sampled: 03/14/07
Received: 03/14/07

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC1612-06 (MWG004_WG031407_0001 - Water) - cont.									
Reporting Units: ug/l									
p-Isopropyltoluene	EPA 8260B	7C16022	0.28	1.0	ND	1	03/16/07	03/16/07	
Methyl-tert-butyl Ether (MTBE)	EPA 8260B	7C16022	0.32	1.0	ND	1	03/16/07	03/16/07	
Methylene chloride	EPA 8260B	7C16022	0.95	1.0	ND	1	03/16/07	03/16/07	
4-Methyl-2-pentanone (MIBK)	EPA 8260B	7C16022	3.5	5.0	ND	1	03/16/07	03/16/07	
n-Propylbenzene	EPA 8260B	7C16022	0.27	1.0	ND	1	03/16/07	03/16/07	
Styrene	EPA 8260B	7C16022	0.16	1.0	ND	1	03/16/07	03/16/07	
1,1,1,2-Tetrachloroethane	EPA 8260B	7C16022	0.27	1.0	ND	1	03/16/07	03/16/07	
1,1,2,2-Tetrachloroethane	EPA 8260B	7C16022	0.24	1.0	ND	1	03/16/07	03/16/07	
Tetrachloroethene	EPA 8260B	7C16022	0.32	1.0	ND	1	03/16/07	03/16/07	
Tetrahydrofuran (THF)	EPA 8260B	7C16022	3.5	10	ND	1	03/16/07	03/16/07	
Toluene	EPA 8260B	7C16022	0.36	1.0	ND	1	03/16/07	03/16/07	
1,2,3-Trichlorobenzene	EPA 8260B	7C16022	0.30	1.0	ND	1	03/16/07	03/16/07	
1,2,4-Trichlorobenzene	EPA 8260B	7C16022	0.48	1.0	ND	1	03/16/07	03/16/07	
1,1,2-Trichloroethane	EPA 8260B	7C16022	0.30	1.0	ND	1	03/16/07	03/16/07	
1,1,1-Trichloroethane	EPA 8260B	7C16022	0.30	1.0	ND	1	03/16/07	03/16/07	
Trichloroethene	EPA 8260B	7C16022	0.26	1.0	17	1	03/16/07	03/16/07	
Trichlorofluoromethane	EPA 8260B	7C16022	0.34	2.0	ND	1	03/16/07	03/16/07	
1,2,3-Trichloropropane	EPA 8260B	7C16022	0.40	1.0	ND	1	03/16/07	03/16/07	
1,2,4-Trimethylbenzene	EPA 8260B	7C16022	0.23	1.0	ND	1	03/16/07	03/16/07	
1,3,5-Trimethylbenzene	EPA 8260B	7C16022	0.26	1.0	ND	1	03/16/07	03/16/07	
Vinyl acetate	EPA 8260B	7C16022	1.7	6.0	ND	1	03/16/07	03/16/07	
Vinyl chloride	EPA 8260B	7C16022	0.30	0.50	ND	1	03/16/07	03/16/07	
Xylenes, Total	EPA 8260B	7C16022	0.90	1.0	ND	1	03/16/07	03/16/07	
Surrogate: 4-Bromofluorobenzene (80-120%)									107 %
Surrogate: Dibromofluoromethane (80-120%)									114 %
Surrogate: Toluene-d8 (80-120%)									104 %

TestAmerica - Irvine, CA
Nicholas Marz
Project Manager

03/15/07

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TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727-01
Report Number: IQC1612

Sampled: 03/14/07
Received: 03/14/07

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC1612-07 (TMW_14_WG031407_0001 - Water)									
Reporting Units: ug/l									
Acetone	EPA 8260B	7C16022	4.5	10	ND	1	03/16/07	03/17/07	
Benzene	EPA 8260B	7C16022	0.28	1.0	ND	1	03/16/07	03/17/07	
Bromobenzene	EPA 8260B	7C16022	0.27	1.0	ND	1	03/16/07	03/17/07	
Bromochloromethane	EPA 8260B	7C16022	0.32	1.0	ND	1	03/16/07	03/17/07	
Bromodichloromethane	EPA 8260B	7C16022	0.30	1.0	ND	1	03/16/07	03/17/07	
Bromoform	EPA 8260B	7C16022	0.40	1.0	ND	1	03/16/07	03/17/07	
Bromomethane	EPA 8260B	7C16022	0.42	1.0	ND	1	03/16/07	03/17/07	
2-Butanone (MEK)	EPA 8260B	7C16022	3.8	5.0	ND	1	03/16/07	03/17/07	
n-Butylbenzene	EPA 8260B	7C16022	0.37	1.0	ND	1	03/16/07	03/17/07	
sec-Butylbenzene	EPA 8260B	7C16022	0.25	1.0	ND	1	03/16/07	03/17/07	
tert-Butylbenzene	EPA 8260B	7C16022	0.22	1.0	ND	1	03/16/07	03/17/07	
Carbon Disulfide	EPA 8260B	7C16022	0.48	1.0	ND	1	03/16/07	03/17/07	
Carbon tetrachloride	EPA 8260B	7C16022	0.28	0.50	1.4	1	03/16/07	03/17/07	
Chlorobenzene	EPA 8260B	7C16022	0.36	1.0	ND	1	03/16/07	03/17/07	
Chloroethane	EPA 8260B	7C16022	0.40	2.0	ND	1	03/16/07	03/17/07	
Chloroform	EPA 8260B	7C16022	0.33	1.0	2.7	1	03/16/07	03/17/07	
Chloromethane	EPA 8260B	7C16022	0.40	2.0	ND	1	03/16/07	03/17/07	
2-Chlorotoluene	EPA 8260B	7C16022	0.28	1.0	ND	1	03/16/07	03/17/07	
4-Chlorotoluene	EPA 8260B	7C16022	0.29	1.0	ND	1	03/16/07	03/17/07	
1,2-Dibromo-3-chloropropane	EPA 8260B	7C16022	0.97	2.0	ND	1	03/16/07	03/17/07	
Dibromochloromethane	EPA 8260B	7C16022	0.28	1.0	ND	1	03/16/07	03/17/07	
1,2-Dibromoethane (EDB)	EPA 8260B	7C16022	0.40	1.0	ND	1	03/16/07	03/17/07	
1,4-Dichlorobenzene	EPA 8260B	7C16022	0.37	1.0	ND	1	03/16/07	03/17/07	
1,2-Dichlorobenzene	EPA 8260B	7C16022	0.32	1.0	ND	1	03/16/07	03/17/07	
1,3-Dichlorobenzene	EPA 8260B	7C16022	0.35	1.0	ND	1	03/16/07	03/17/07	
Dichlorodifluoromethane	EPA 8260B	7C16022	0.79	1.0	ND	1	03/16/07	03/17/07	
1,2-Dichloroethane	EPA 8260B	7C16022	0.28	0.50	ND	1	03/16/07	03/17/07	
1,1-Dichloroethane	EPA 8260B	7C16022	0.27	1.0	ND	1	03/16/07	03/17/07	
1,1-Dichloroethene	EPA 8260B	7C16022	0.42	1.0	ND	1	03/16/07	03/17/07	
cis-1,2-Dichloroethene	EPA 8260B	7C16022	0.32	1.0	ND	1	03/16/07	03/17/07	
trans-1,2-Dichloroethene	EPA 8260B	7C16022	0.27	1.0	ND	1	03/16/07	03/17/07	
1,2-Dichloropropane	EPA 8260B	7C16022	0.35	1.0	ND	1	03/16/07	03/17/07	
2,2-Dichloropropane	EPA 8260B	7C16022	0.34	1.0	ND	1	03/16/07	03/17/07	
cis-1,3-Dichloropropene	EPA 8260B	7C16022	0.22	0.50	ND	1	03/16/07	03/17/07	
1,1-Dichloropropene	EPA 8260B	7C16022	0.28	1.0	ND	1	03/16/07	03/17/07	
trans-1,3-Dichloropropene	EPA 8260B	7C16022	0.32	0.50	ND	1	03/16/07	03/17/07	
Ethylbenzene	EPA 8260B	7C16022	0.25	1.0	ND	1	03/16/07	03/17/07	
Hexachlorobutadiene	EPA 8260B	7C16022	0.38	1.0	ND	1	03/16/07	03/17/07	
2-Hexanone	EPA 8260B	7C16022	2.6	6.0	ND	1	03/16/07	03/17/07	
Iodomethane	EPA 8260B	7C16022	1.0	2.0	ND	1	03/16/07	03/17/07	
Isopropylbenzene	EPA 8260B	7C16022	0.25	1.0	ND	1	03/16/07	03/17/07	

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Nicholas Marz
Project Manager

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ANALYTICAL TESTING CORPORATION

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TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727-01
Report Number: IQC1612

Sampled: 03/14/07
Received: 03/14/07

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC1612-07 (TMW_14_WG031407_0001 - Water) - cont.									
Reporting Units: ug/l									
p-Isopropyltoluene	EPA 8260B	7C16022	0.28	1.0	ND	1	03/16/07	03/17/07	
Methyl-tert-butyl Ether (MTBE)	EPA 8260B	7C16022	0.32	1.0	ND	1	03/16/07	03/17/07	
Methylene chloride	EPA 8260B	7C16022	0.95	1.0	ND	1	03/16/07	03/17/07	
4-Methyl-2-pentanone (MIBK)	EPA 8260B	7C16022	3.5	5.0	ND	1	03/16/07	03/17/07	
n-Propylbenzene	EPA 8260B	7C16022	0.27	1.0	ND	1	03/16/07	03/17/07	
Styrene	EPA 8260B	7C16022	0.16	1.0	ND	1	03/16/07	03/17/07	
1,1,1,2-Tetrachloroethane	EPA 8260B	7C16022	0.27	1.0	ND	1	03/16/07	03/17/07	
1,1,2,2-Tetrachloroethane	EPA 8260B	7C16022	0.24	1.0	ND	1	03/16/07	03/17/07	
Tetrachloroethene	EPA 8260B	7C16022	0.32	1.0	0.89	1	03/16/07	03/17/07	J
Tetrahydrofuran (THF)	EPA 8260B	7C16022	3.5	10	ND	1	03/16/07	03/17/07	
Toluene	EPA 8260B	7C16022	0.36	1.0	ND	1	03/16/07	03/17/07	
1,2,3-Trichlorobenzene	EPA 8260B	7C16022	0.30	1.0	ND	1	03/16/07	03/17/07	
1,2,4-Trichlorobenzene	EPA 8260B	7C16022	0.48	1.0	ND	1	03/16/07	03/17/07	
1,1,2-Trichloroethane	EPA 8260B	7C16022	0.30	1.0	ND	1	03/16/07	03/17/07	
1,1,1-Trichloroethane	EPA 8260B	7C16022	0.30	1.0	ND	1	03/16/07	03/17/07	
Trichloroethene	EPA 8260B	7C16022	0.26	1.0	6.7	1	03/16/07	03/17/07	
Trichlorofluoromethane	EPA 8260B	7C16022	0.34	2.0	ND	1	03/16/07	03/17/07	
1,2,3-Trichloropropane	EPA 8260B	7C16022	0.40	1.0	ND	1	03/16/07	03/17/07	
1,2,4-Trimethylbenzene	EPA 8260B	7C16022	0.23	1.0	ND	1	03/16/07	03/17/07	
1,3,5-Trimethylbenzene	EPA 8260B	7C16022	0.26	1.0	ND	1	03/16/07	03/17/07	
Vinyl acetate	EPA 8260B	7C16022	1.7	6.0	ND	1	03/16/07	03/17/07	
Vinyl chloride	EPA 8260B	7C16022	0.30	0.50	ND	1	03/16/07	03/17/07	
Xylenes, Total	EPA 8260B	7C16022	0.90	1.0	ND	1	03/16/07	03/17/07	
Surrogate: 4-Bromofluorobenzene (80-120%)									108 %
Surrogate: Dibromofluoromethane (80-120%)									118 %
Surrogate: Toluene-d8 (80-120%)									104 %

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Nicholas Marz
Project Manager

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ANALYTICAL TESTING CORPORATION

17461 Derian Avenue, Suite 100, Irvine, CA 92614 (949) 261-1022 Fax: (949) 260-3297

TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727-01
Report Number: IQC1612

Sampled: 03/14/07
Received: 03/14/07

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC1612-08 (TMW_11_WG031407_0001 - Water)									
Reporting Units: ug/l									
Acetone	EPA 8260B	7C16022	4.5	10	ND	1	03/16/07	03/17/07	
Benzene	EPA 8260B	7C16022	0.28	1.0	0.28	1	03/16/07	03/17/07	J
Bromobenzene	EPA 8260B	7C16022	0.27	1.0	ND	1	03/16/07	03/17/07	
Bromochloromethane	EPA 8260B	7C16022	0.32	1.0	ND	1	03/16/07	03/17/07	
Bromodichloromethane	EPA 8260B	7C16022	0.30	1.0	ND	1	03/16/07	03/17/07	
Bromoform	EPA 8260B	7C16022	0.40	1.0	ND	1	03/16/07	03/17/07	
Bromomethane	EPA 8260B	7C16022	0.42	1.0	ND	1	03/16/07	03/17/07	
2-Butanone (MEK)	EPA 8260B	7C16022	3.8	5.0	ND	1	03/16/07	03/17/07	
n-Butylbenzene	EPA 8260B	7C16022	0.37	1.0	ND	1	03/16/07	03/17/07	
sec-Butylbenzene	EPA 8260B	7C16022	0.25	1.0	ND	1	03/16/07	03/17/07	
tert-Butylbenzene	EPA 8260B	7C16022	0.22	1.0	ND	1	03/16/07	03/17/07	
Carbon Disulfide	EPA 8260B	7C16022	0.48	1.0	ND	1	03/16/07	03/17/07	
Carbon tetrachloride	EPA 8260B	7C16022	0.28	0.50	2.3	1	03/16/07	03/17/07	
Chlorobenzene	EPA 8260B	7C16022	0.36	1.0	ND	1	03/16/07	03/17/07	
Chloroethane	EPA 8260B	7C16022	0.40	2.0	ND	1	03/16/07	03/17/07	
Chloroform	EPA 8260B	7C16022	0.33	1.0	170	1	03/16/07	03/17/07	
Chloromethane	EPA 8260B	7C16022	0.40	2.0	ND	1	03/16/07	03/17/07	
2-Chlorotoluene	EPA 8260B	7C16022	0.28	1.0	ND	1	03/16/07	03/17/07	
4-Chlorotoluene	EPA 8260B	7C16022	0.29	1.0	ND	1	03/16/07	03/17/07	
1,2-Dibromo-3-chloropropane	EPA 8260B	7C16022	0.97	2.0	ND	1	03/16/07	03/17/07	
Dibromochloromethane	EPA 8260B	7C16022	0.28	1.0	ND	1	03/16/07	03/17/07	
1,2-Dibromoethane (EDB)	EPA 8260B	7C16022	0.40	1.0	ND	1	03/16/07	03/17/07	
1,4-Dichlorobenzene	EPA 8260B	7C16022	0.37	1.0	ND	1	03/16/07	03/17/07	
1,2-Dichlorobenzene	EPA 8260B	7C16022	0.32	1.0	ND	1	03/16/07	03/17/07	
1,3-Dichlorobenzene	EPA 8260B	7C16022	0.35	1.0	ND	1	03/16/07	03/17/07	
Dichlorodifluoromethane	EPA 8260B	7C16022	0.79	1.0	ND	1	03/16/07	03/17/07	
1,2-Dichloroethane	EPA 8260B	7C16022	0.28	0.50	ND	1	03/16/07	03/17/07	
1,1-Dichloroethane	EPA 8260B	7C16022	0.27	1.0	ND	1	03/16/07	03/17/07	
1,1-Dichloroethene	EPA 8260B	7C16022	0.42	1.0	ND	1	03/16/07	03/17/07	
cis-1,2-Dichloroethene	EPA 8260B	7C16022	0.32	1.0	ND	1	03/16/07	03/17/07	
trans-1,2-Dichloroethene	EPA 8260B	7C16022	0.27	1.0	ND	1	03/16/07	03/17/07	
1,2-Dichloropropane	EPA 8260B	7C16022	0.35	1.0	ND	1	03/16/07	03/17/07	
2,2-Dichloropropane	EPA 8260B	7C16022	0.34	1.0	ND	1	03/16/07	03/17/07	
cis-1,3-Dichloropropene	EPA 8260B	7C16022	0.22	0.50	ND	1	03/16/07	03/17/07	
1,1-Dichloropropene	EPA 8260B	7C16022	0.28	1.0	ND	1	03/16/07	03/17/07	
trans-1,3-Dichloropropene	EPA 8260B	7C16022	0.32	0.50	ND	1	03/16/07	03/17/07	
Ethylbenzene	EPA 8260B	7C16022	0.25	1.0	ND	1	03/16/07	03/17/07	
Hexachlorobutadiene	EPA 8260B	7C16022	0.38	1.0	ND	1	03/16/07	03/17/07	
2-Hexanone	EPA 8260B	7C16022	2.6	6.0	ND	1	03/16/07	03/17/07	
Iodomethane	EPA 8260B	7C16022	1.0	2.0	ND	1	03/16/07	03/17/07	
Isopropylbenzene	EPA 8260B	7C16022	0.25	1.0	ND	1	03/16/07	03/17/07	

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Nicholas Marz
Project Manager

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ANALYTICAL TESTING CORPORATION

17461 Derian Avenue, Suite 100, Irvine, CA 92614 (949) 261-1022 Fax: (949) 260-3297

TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727-01
Report Number: IQC1612

Sampled: 03/14/07
Received: 03/14/07

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC1612-08 (TMW_11_WG031407_0001 - Water) - cont.									
Reporting Units: ug/l									
p-Isopropyltoluene	EPA 8260B	7C16022	0.28	1.0	ND	1	03/16/07	03/17/07	
Methyl-tert-butyl Ether (MTBE)	EPA 8260B	7C16022	0.32	1.0	ND	1	03/16/07	03/17/07	
Methylene chloride	EPA 8260B	7C16022	0.95	1.0	ND	1	03/16/07	03/17/07	
4-Methyl-2-pentanone (MIBK)	EPA 8260B	7C16022	3.5	5.0	ND	1	03/16/07	03/17/07	
n-Propylbenzene	EPA 8260B	7C16022	0.27	1.0	ND	1	03/16/07	03/17/07	
Styrene	EPA 8260B	7C16022	0.16	1.0	ND	1	03/16/07	03/17/07	
1,1,1,2-Tetrachloroethane	EPA 8260B	7C16022	0.27	1.0	ND	1	03/16/07	03/17/07	
1,1,2,2-Tetrachloroethane	EPA 8260B	7C16022	0.24	1.0	ND	1	03/16/07	03/17/07	
Tetrachloroethene	EPA 8260B	7C16022	0.32	1.0	3.7	1	03/16/07	03/17/07	
Tetrahydrofuran (THF)	EPA 8260B	7C16022	3.5	10	ND	1	03/16/07	03/17/07	
Toluene	EPA 8260B	7C16022	0.36	1.0	ND	1	03/16/07	03/17/07	
1,2,3-Trichlorobenzene	EPA 8260B	7C16022	0.30	1.0	ND	1	03/16/07	03/17/07	
1,2,4-Trichlorobenzene	EPA 8260B	7C16022	0.48	1.0	ND	1	03/16/07	03/17/07	
1,1,2-Trichloroethane	EPA 8260B	7C16022	0.30	1.0	ND	1	03/16/07	03/17/07	
1,1,1-Trichloroethane	EPA 8260B	7C16022	0.30	1.0	ND	1	03/16/07	03/17/07	
Trichloroethene	EPA 8260B	7C16022	0.26	1.0	8.0	1	03/16/07	03/17/07	
Trichlorofluoromethane	EPA 8260B	7C16022	0.34	2.0	ND	1	03/16/07	03/17/07	
1,2,3-Trichloropropane	EPA 8260B	7C16022	0.40	1.0	ND	1	03/16/07	03/17/07	
1,2,4-Trimethylbenzene	EPA 8260B	7C16022	0.23	1.0	ND	1	03/16/07	03/17/07	
1,3,5-Trimethylbenzene	EPA 8260B	7C16022	0.26	1.0	ND	1	03/16/07	03/17/07	
Vinyl acetate	EPA 8260B	7C16022	1.7	6.0	ND	1	03/16/07	03/17/07	
Vinyl chloride	EPA 8260B	7C16022	0.30	0.50	ND	1	03/16/07	03/17/07	
Xylenes, Total	EPA 8260B	7C16022	0.90	1.0	ND	1	03/16/07	03/17/07	
Surrogate: 4-Bromofluorobenzene (80-120%)									109 %
Surrogate: Dibromofluoromethane (80-120%)									111 %
Surrogate: Toluene-d8 (80-120%)									102 %

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Nicholas Marz
Project Manager

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ANALYTICAL TESTING CORPORATION

17461 Derian Avenue, Suite 100, Irvine, CA 92614 (949) 261-1022 Fax: (949) 260-3297

TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727-01
Report Number: IQC1612

Sampled: 03/14/07
Received: 03/14/07

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC1612-09 (WCC_5S_WG031407_0001 - Water)									
Reporting Units: ug/l									
Acetone	EPA 8260B	7C16022	4.5	10	ND	1	03/16/07	03/17/07	
Benzene	EPA 8260B	7C16022	0.28	1.0	ND	1	03/16/07	03/17/07	
Bromobenzene	EPA 8260B	7C16022	0.27	1.0	ND	1	03/16/07	03/17/07	
Bromochloromethane	EPA 8260B	7C16022	0.32	1.0	ND	1	03/16/07	03/17/07	
Bromodichloromethane	EPA 8260B	7C16022	0.30	1.0	ND	1	03/16/07	03/17/07	
Bromoform	EPA 8260B	7C16022	0.40	1.0	ND	1	03/16/07	03/17/07	
Bromomethane	EPA 8260B	7C16022	0.42	1.0	ND	1	03/16/07	03/17/07	
2-Butanone (MEK)	EPA 8260B	7C16022	3.8	5.0	ND	1	03/16/07	03/17/07	
n-Butylbenzene	EPA 8260B	7C16022	0.37	1.0	ND	1	03/16/07	03/17/07	
sec-Butylbenzene	EPA 8260B	7C16022	0.25	1.0	ND	1	03/16/07	03/17/07	
tert-Butylbenzene	EPA 8260B	7C16022	0.22	1.0	ND	1	03/16/07	03/17/07	
Carbon Disulfide	EPA 8260B	7C16022	0.48	1.0	ND	1	03/16/07	03/17/07	
Carbon tetrachloride	EPA 8260B	7C16022	0.28	0.50	ND	1	03/16/07	03/17/07	
Chlorobenzene	EPA 8260B	7C16022	0.36	1.0	ND	1	03/16/07	03/17/07	
Chloroethane	EPA 8260B	7C16022	0.40	2.0	ND	1	03/16/07	03/17/07	
Chloroform	EPA 8260B	7C16022	0.33	1.0	0.40	1	03/16/07	03/17/07	J
Chloromethane	EPA 8260B	7C16022	0.40	2.0	ND	1	03/16/07	03/17/07	
2-Chlorotoluene	EPA 8260B	7C16022	0.28	1.0	ND	1	03/16/07	03/17/07	
4-Chlorotoluene	EPA 8260B	7C16022	0.29	1.0	ND	1	03/16/07	03/17/07	
1,2-Dibromo-3-chloropropane	EPA 8260B	7C16022	0.97	2.0	ND	1	03/16/07	03/17/07	
Dibromochloromethane	EPA 8260B	7C16022	0.28	1.0	ND	1	03/16/07	03/17/07	
1,2-Dibromoethane (EDB)	EPA 8260B	7C16022	0.40	1.0	ND	1	03/16/07	03/17/07	
1,4-Dichlorobenzene	EPA 8260B	7C16022	0.37	1.0	ND	1	03/16/07	03/17/07	
1,2-Dichlorobenzene	EPA 8260B	7C16022	0.32	1.0	ND	1	03/16/07	03/17/07	
1,3-Dichlorobenzene	EPA 8260B	7C16022	0.35	1.0	ND	1	03/16/07	03/17/07	
Dichlorodifluoromethane	EPA 8260B	7C16022	0.79	1.0	ND	1	03/16/07	03/17/07	
1,2-Dichloroethane	EPA 8260B	7C16022	0.28	0.50	ND	1	03/16/07	03/17/07	
1,1-Dichloroethane	EPA 8260B	7C16022	0.27	1.0	0.68	1	03/16/07	03/17/07	J
1,1-Dichloroethene	EPA 8260B	7C16022	0.42	1.0	6.5	1	03/16/07	03/17/07	
cis-1,2-Dichloroethene	EPA 8260B	7C16022	0.32	1.0	0.54	1	03/16/07	03/17/07	J
trans-1,2-Dichloroethene	EPA 8260B	7C16022	0.27	1.0	ND	1	03/16/07	03/17/07	
1,2-Dichloropropane	EPA 8260B	7C16022	0.35	1.0	ND	1	03/16/07	03/17/07	
2,2-Dichloropropane	EPA 8260B	7C16022	0.34	1.0	ND	1	03/16/07	03/17/07	
cis-1,3-Dichloropropene	EPA 8260B	7C16022	0.22	0.50	ND	1	03/16/07	03/17/07	
1,1-Dichloropropene	EPA 8260B	7C16022	0.28	1.0	ND	1	03/16/07	03/17/07	
trans-1,3-Dichloropropene	EPA 8260B	7C16022	0.32	0.50	ND	1	03/16/07	03/17/07	
Ethylbenzene	EPA 8260B	7C16022	0.25	1.0	ND	1	03/16/07	03/17/07	
Hexachlorobutadiene	EPA 8260B	7C16022	0.38	1.0	ND	1	03/16/07	03/17/07	
2-Hexanone	EPA 8260B	7C16022	2.6	6.0	ND	1	03/16/07	03/17/07	
Iodomethane	EPA 8260B	7C16022	1.0	2.0	ND	1	03/16/07	03/17/07	
Isopropylbenzene	EPA 8260B	7C16022	0.25	1.0	ND	1	03/16/07	03/17/07	

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Project Manager

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TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727-01
Report Number: IQC1612

Sampled: 03/14/07
Received: 03/14/07

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC1612-09 (WCC_5S_WG031407_0001 - Water) - cont.									
Reporting Units: ug/l									
p-Isopropyltoluene	EPA 8260B	7C16022	0.28	1.0	ND	1	03/16/07	03/17/07	
Methyl-tert-butyl Ether (MTBE)	EPA 8260B	7C16022	0.32	1.0	ND	1	03/16/07	03/17/07	
Methylene chloride	EPA 8260B	7C16022	0.95	1.0	ND	1	03/16/07	03/17/07	
4-Methyl-2-pentanone (MIBK)	EPA 8260B	7C16022	3.5	5.0	ND	1	03/16/07	03/17/07	
n-Propylbenzene	EPA 8260B	7C16022	0.27	1.0	ND	1	03/16/07	03/17/07	
Styrene	EPA 8260B	7C16022	0.16	1.0	ND	1	03/16/07	03/17/07	
1,1,1,2-Tetrachloroethane	EPA 8260B	7C16022	0.27	1.0	ND	1	03/16/07	03/17/07	
1,1,2,2-Tetrachloroethane	EPA 8260B	7C16022	0.24	1.0	ND	1	03/16/07	03/17/07	
Tetrachloroethene	EPA 8260B	7C16022	0.32	1.0	ND	1	03/16/07	03/17/07	
Tetrahydrofuran (THF)	EPA 8260B	7C16022	3.5	10	ND	1	03/16/07	03/17/07	
Toluene	EPA 8260B	7C16022	0.36	1.0	ND	1	03/16/07	03/17/07	
1,2,3-Trichlorobenzene	EPA 8260B	7C16022	0.30	1.0	ND	1	03/16/07	03/17/07	
1,2,4-Trichlorobenzene	EPA 8260B	7C16022	0.48	1.0	ND	1	03/16/07	03/17/07	
1,1,2-Trichloroethane	EPA 8260B	7C16022	0.30	1.0	ND	1	03/16/07	03/17/07	
1,1,1-Trichloroethane	EPA 8260B	7C16022	0.30	1.0	ND	1	03/16/07	03/17/07	
Trichloroethene	EPA 8260B	7C16022	0.26	1.0	2.9	1	03/16/07	03/17/07	
Trichlorofluoromethane	EPA 8260B	7C16022	0.34	2.0	1.2	1	03/16/07	03/17/07	J
1,2,3-Trichloropropane	EPA 8260B	7C16022	0.40	1.0	ND	1	03/16/07	03/17/07	
1,2,4-Trimethylbenzene	EPA 8260B	7C16022	0.23	1.0	ND	1	03/16/07	03/17/07	
1,3,5-Trimethylbenzene	EPA 8260B	7C16022	0.26	1.0	ND	1	03/16/07	03/17/07	
Vinyl acetate	EPA 8260B	7C16022	1.7	6.0	ND	1	03/16/07	03/17/07	
Vinyl chloride	EPA 8260B	7C16022	0.30	0.50	ND	1	03/16/07	03/17/07	
Xylenes, Total	EPA 8260B	7C16022	0.90	1.0	ND	1	03/16/07	03/17/07	
Surrogate: 4-Bromofluorobenzene (80-120%)									109 %
Surrogate: Dibromofluoromethane (80-120%)									116 %
Surrogate: Toluene-d8 (80-120%)									103 %

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Nicholas Marz
Project Manager

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ANALYTICAL TESTING CORPORATION

17461 Derian Avenue, Suite 100, Irvine, CA 92614 (949) 261-1022 Fax: (949) 260-3297

TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727-01
Report Number: IQC1612

Sampled: 03/14/07
Received: 03/14/07

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC1612-10 (MWC021_WG031407_0001 - Water)									
Reporting Units: ug/l									
Acetone	EPA 8260B	7C16022	4.5	10	ND	1	03/16/07	03/17/07	
Benzene	EPA 8260B	7C16022	0.28	1.0	ND	1	03/16/07	03/17/07	
Bromobenzene	EPA 8260B	7C16022	0.27	1.0	ND	1	03/16/07	03/17/07	
Bromochloromethane	EPA 8260B	7C16022	0.32	1.0	ND	1	03/16/07	03/17/07	
Bromodichloromethane	EPA 8260B	7C16022	0.30	1.0	ND	1	03/16/07	03/17/07	
Bromoform	EPA 8260B	7C16022	0.40	1.0	ND	1	03/16/07	03/17/07	
Bromomethane	EPA 8260B	7C16022	0.42	1.0	ND	1	03/16/07	03/17/07	
2-Butanone (MEK)	EPA 8260B	7C16022	3.8	5.0	ND	1	03/16/07	03/17/07	
n-Butylbenzene	EPA 8260B	7C16022	0.37	1.0	ND	1	03/16/07	03/17/07	
sec-Butylbenzene	EPA 8260B	7C16022	0.25	1.0	ND	1	03/16/07	03/17/07	
tert-Butylbenzene	EPA 8260B	7C16022	0.22	1.0	ND	1	03/16/07	03/17/07	
Carbon Disulfide	EPA 8260B	7C16022	0.48	1.0	ND	1	03/16/07	03/17/07	
Carbon tetrachloride	EPA 8260B	7C16022	0.28	0.50	ND	1	03/16/07	03/17/07	
Chlorobenzene	EPA 8260B	7C16022	0.36	1.0	ND	1	03/16/07	03/17/07	
Chloroethane	EPA 8260B	7C16022	0.40	2.0	ND	1	03/16/07	03/17/07	
Chloroform	EPA 8260B	7C16022	0.33	1.0	0.50	1	03/16/07	03/17/07	J
Chloromethane	EPA 8260B	7C16022	0.40	2.0	ND	1	03/16/07	03/17/07	
2-Chlorotoluene	EPA 8260B	7C16022	0.28	1.0	ND	1	03/16/07	03/17/07	
4-Chlorotoluene	EPA 8260B	7C16022	0.29	1.0	ND	1	03/16/07	03/17/07	
1,2-Dibromo-3-chloropropane	EPA 8260B	7C16022	0.97	2.0	ND	1	03/16/07	03/17/07	
Dibromochloromethane	EPA 8260B	7C16022	0.28	1.0	ND	1	03/16/07	03/17/07	
1,2-Dibromoethane (EDB)	EPA 8260B	7C16022	0.40	1.0	ND	1	03/16/07	03/17/07	
1,4-Dichlorobenzene	EPA 8260B	7C16022	0.37	1.0	ND	1	03/16/07	03/17/07	
1,2-Dichlorobenzene	EPA 8260B	7C16022	0.32	1.0	ND	1	03/16/07	03/17/07	
1,3-Dichlorobenzene	EPA 8260B	7C16022	0.35	1.0	ND	1	03/16/07	03/17/07	
Dichlorodifluoromethane	EPA 8260B	7C16022	0.79	1.0	ND	1	03/16/07	03/17/07	
1,2-Dichloroethane	EPA 8260B	7C16022	0.28	0.50	ND	1	03/16/07	03/17/07	
1,1-Dichloroethane	EPA 8260B	7C16022	0.27	1.0	0.52	1	03/16/07	03/17/07	J
1,1-Dichloroethene	EPA 8260B	7C16022	0.42	1.0	0.46	1	03/16/07	03/17/07	J
cis-1,2-Dichloroethene	EPA 8260B	7C16022	0.32	1.0	4.1	1	03/16/07	03/17/07	
trans-1,2-Dichloroethene	EPA 8260B	7C16022	0.27	1.0	ND	1	03/16/07	03/17/07	
1,2-Dichloropropane	EPA 8260B	7C16022	0.35	1.0	ND	1	03/16/07	03/17/07	
2,2-Dichloropropane	EPA 8260B	7C16022	0.34	1.0	ND	1	03/16/07	03/17/07	
cis-1,3-Dichloropropene	EPA 8260B	7C16022	0.22	0.50	ND	1	03/16/07	03/17/07	
1,1-Dichloropropene	EPA 8260B	7C16022	0.28	1.0	ND	1	03/16/07	03/17/07	
trans-1,3-Dichloropropene	EPA 8260B	7C16022	0.32	0.50	ND	1	03/16/07	03/17/07	
Ethylbenzene	EPA 8260B	7C16022	0.25	1.0	ND	1	03/16/07	03/17/07	
Hexachlorobutadiene	EPA 8260B	7C16022	0.38	1.0	ND	1	03/16/07	03/17/07	
2-Hexanone	EPA 8260B	7C16022	2.6	6.0	ND	1	03/16/07	03/17/07	
Iodomethane	EPA 8260B	7C16022	1.0	2.0	ND	1	03/16/07	03/17/07	
Isopropylbenzene	EPA 8260B	7C16022	0.25	1.0	ND	1	03/16/07	03/17/07	

TestAmerica - Irvine, CA
Nicholas Marz
Project Manager

RAC1007

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TestAmerica

ANALYTICAL TESTING CORPORATION

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TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727-01
Report Number: IQC1612

Sampled: 03/14/07
Received: 03/14/07

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC1612-10 (MWC021_WG031407_0001 - Water) - cont.									
Reporting Units: ug/l									
p-Isopropyltoluene	EPA 8260B	7C16022	0.28	1.0	ND	1	03/16/07	03/17/07	
Methyl-tert-butyl Ether (MTBE)	EPA 8260B	7C16022	0.32	1.0	ND	1	03/16/07	03/17/07	
Methylene chloride	EPA 8260B	7C16022	0.95	1.0	ND	1	03/16/07	03/17/07	
4-Methyl-2-pentanone (MIBK)	EPA 8260B	7C16022	3.5	5.0	ND	1	03/16/07	03/17/07	
n-Propylbenzene	EPA 8260B	7C16022	0.27	1.0	ND	1	03/16/07	03/17/07	
Styrene	EPA 8260B	7C16022	0.16	1.0	ND	1	03/16/07	03/17/07	
1,1,1,2-Tetrachloroethane	EPA 8260B	7C16022	0.27	1.0	ND	1	03/16/07	03/17/07	
1,1,2,2-Tetrachloroethane	EPA 8260B	7C16022	0.24	1.0	ND	1	03/16/07	03/17/07	
Tetrachloroethene	EPA 8260B	7C16022	0.32	1.0	ND	1	03/16/07	03/17/07	
Tetrahydrofuran (THF)	EPA 8260B	7C16022	3.5	10	ND	1	03/16/07	03/17/07	
Toluene	EPA 8260B	7C16022	0.36	1.0	ND	1	03/16/07	03/17/07	
1,2,3-Trichlorobenzene	EPA 8260B	7C16022	0.30	1.0	ND	1	03/16/07	03/17/07	
1,2,4-Trichlorobenzene	EPA 8260B	7C16022	0.48	1.0	ND	1	03/16/07	03/17/07	
1,1,2-Trichloroethane	EPA 8260B	7C16022	0.30	1.0	ND	1	03/16/07	03/17/07	
1,1,1-Trichloroethane	EPA 8260B	7C16022	0.30	1.0	ND	1	03/16/07	03/17/07	
Trichloroethene	EPA 8260B	7C16022	0.26	1.0	11	1	03/16/07	03/17/07	
Trichlorofluoromethane	EPA 8260B	7C16022	0.34	2.0	ND	1	03/16/07	03/17/07	
1,2,3-Trichloropropane	EPA 8260B	7C16022	0.40	1.0	ND	1	03/16/07	03/17/07	
1,2,4-Trimethylbenzene	EPA 8260B	7C16022	0.23	1.0	ND	1	03/16/07	03/17/07	
1,3,5-Trimethylbenzene	EPA 8260B	7C16022	0.26	1.0	ND	1	03/16/07	03/17/07	
Vinyl acetate	EPA 8260B	7C16022	1.7	6.0	ND	1	03/16/07	03/17/07	
Vinyl chloride	EPA 8260B	7C16022	0.30	0.50	ND	1	03/16/07	03/17/07	
Xylenes, Total	EPA 8260B	7C16022	0.90	1.0	ND	1	03/16/07	03/17/07	
Surrogate: 4-Bromofluorobenzene (80-120%)					109 %				
Surrogate: Dibromofluoromethane (80-120%)					115 %				
Surrogate: Toluene-d8 (80-120%)					104 %				

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Nicholas Marz
Project Manager

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LDC #: 16739A1
 SDG #: IQC1612
 Laboratory: Test America

VALIDATION COMPLETENESS WORKSHEET
 Tier 1

Date: 5/10/07
 Page: 1 of 1
 Reviewer: P
 2nd Reviewer: R

METHOD: GC/MS Volatiles (EPA SW 846 Method 8260B)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: 3/14/07
II.	GC/MS Instrument performance check	N	
III.	Initial calibration	N	
IV.	Continuing calibration	N	
V.	Blanks	A	
VI.	Surrogate spikes	A	
VII.	Matrix spike/Matrix spike duplicates	A	
VIII.	Laboratory control samples	A	
IX.	Regional Quality Assurance and Quality Control	N	
X.	Internal standards	N	
XI.	Target compound identification	N	
XII.	Compound quantitation/CRQLs	N	
XIII.	Tentatively identified compounds (TICs)	N	
XIV.	System performance	N	
XV.	Overall assessment of data	A	
XVI.	Field duplicates	N	
XVII.	Field blanks	N	

Note: A = Acceptable
 N = Not provided/applicable
 SW = See worksheet
 ND = No compounds detected
 R = Rinsate
 FB = Field blank
 D = Duplicate
 TB = Trip blank
 EB = Equipment blank

Validated Samples:
 water

5	1	MWB013_WG031407_0001	11	7c14022 - B41	21	31
6	2	MWG004_WG031407_0001	12		22	32
7	3	TMW_14_WG031407_0001	13		23	33
8	4	TMW_11_WG031407_0001	14		24	34
9	5	WCC_5S_WG031407_0001	15		25	35
10	6	MWC021_WG031407_0001	16		26	36
	7	MWB013_WG031407_0001MS	17		27	37
	8	MWB013_WG031407_0001MSD	18		28	38
	9		19		29	39
	10		20		30	40

LDC Report# 16739B1

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Boeing Realty Corp., Bldg C-6 Torrance
Collection Date: March 22, 2007
LDC Report Date: May 10, 2007
Matrix: Water
Parameters: Volatiles
Validation Level: Tier 2
Laboratory: TestAmerica

Sample Delivery Group (SDG): IQC2470

Sample Identification

MWB028_WG032207_0001
MWB027_WG032207_0001
MWB027_WG032207_0002
CMW002_WG032207_0001
MWB028_WG032207_0001MS
MWB028_WG032207_0001MSD

Introduction

This data review covers 6 water samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 8260B for Volatiles.

This review follows a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Organic Data Review (October 1999) as there are no current guidelines for the method stated above.

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

Blank results are summarized in Section V.

Field duplicates are summarized in Section XVI.

Raw data were not reviewed for this SDG. The review was based on QC data.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- N Presumptive evidence of presence of the constituent.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. GC/MS Instrument Performance Check

Instrument performance was checked at 12 hour intervals.

All ion abundance requirements were met.

III. Initial Calibration

Initial calibration was performed using required standard concentrations.

Percent relative standard deviations (%RSD) were less than or equal to 15.0% for each individual compound and less than or equal to 30.0% for calibration check compounds (CCCs).

In the case where %RSD was greater than 15.0%, the laboratory used a calibration curve to evaluate the compound. All coefficients of determination (r^2) were greater than or equal to 0.990 .

For the purposes of technical evaluation, all compounds were evaluated against the 30.0% (%RSD) National Functional Guideline criteria. Unless noted above, all compounds were within the validation criteria.

Average relative response factors (RRF) for all volatile target compounds and system performance check compounds (SPCCs) were within method and validation criteria with the following exceptions:

Date	Compound	RRF (Limits)	Associated Samples	Flag	A or P
2/27/07	2-Butanone	0.044 (≥ 0.05)	MWB028_WG032207_0001 MWB027_WG032207_0001 MWB027_WG032207_0002 MWB028_WG032207_0001MS MWB028_WG032207_0001MSD 7C27024-BLK1	J (all detects) UJ (all non-detects)	A
3/21/07	2-Butanone	0.047 (≥ 0.05)	CMW002_WG032207_0001 7C29027-BLK1	J (all detects) UJ (all non-detects)	A

IV. Continuing Calibration

Continuing calibration was performed at the required frequencies.

Percent differences (%D) between the initial calibration RRF and the continuing calibration RRF were within the method criteria of less than or equal to 20.0% for calibration check compounds (CCCs).

For the purposes of technical evaluation, all compounds were evaluated against the 25.0% (%D) National Functional Guideline criteria. Unless noted above, all compounds were within the validation criteria with the following exceptions:

Date	Compound	%D	Associated Samples	Flag	A or P
3/27/07	Acetone	27.1	MWB028_WG032207_0001 MWB027_WG032207_0002 MWB028_WG032207_0001MS MWB028_WG032207_0001MSD 7C27024-BLK1	J (all detects) UJ (all non-detects)	A
3/29/07	Tetrahydrofuran 2-Butanone 4-Methyl-2-pentanone 2-Hexanone	44 80.9 26.5 70.8	CMW002_WG032207_0001 7C29027-BLK1	J (all detects) UJ (all non-detects)	A
3/30/07	Acetone	41.3	CMW002_WG032207_0001 7C30007-BLK1	J (all detects) UJ (all non-detects)	A

All of the continuing calibration RRF values were within method and validation criteria.

V. Blanks

Method blanks were reviewed for each matrix as applicable. No volatile contaminants were found in the method blanks with the following exceptions:

Method Blank ID	Analysis Date	Compound TIC (RT in minutes)	Concentration	Associated Samples
7C27024-BLK1	3/27/07	Tetrahydrofuran Toluene	8.04 ug/L 0.440 ug/L	MWB028_WG032207_0001 MWB027_WG032207_0001 MWB027_WG032207_0002
7C29027-BLK1	3/29/07	Tetrahydrofuran	5.88 ug/L	CMW002_WG032207_0001
7C30007-BLK1	3/30/07	Methylene chloride	1.64 ug/L	CMW002_WG032207_0001

Sample concentrations were compared to concentrations detected in the method blanks. The sample concentrations were either not detected or were significantly greater (>10X for common contaminants, >5X for other contaminants) than the concentrations found in the associated method blanks with the following exceptions:

Sample	Compound TIC (RT in minutes)	Reported Concentration	Modified Final Concentration
MWB028_WG032207_0001	Toluene	0.84 ug/L	1.0U ug/L
MWB027_WG032207_0001	Toluene	1.2 ug/L	1.2U ug/L
MWB027_WG032207_0002	Toluene	0.91 ug/L	1.0U ug/L

VI. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VII. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were not within QC limits. Since there were no associated samples, no data were qualified.

VIII. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) were within QC limits with the following exceptions:

LCS ID	Compound	%R (Limits)	Associated Samples	Flag	A or P
7C27024-BS1	Acetone	156 (30-140)	MWB028_WG032207_0001 MWB027_WG032207_0002 7C27024-BLK1	J (all detects)	P
7C27024-BS1	2-Butanone	150 (40-140)	MWB028_WG032207_0001 MWB027_WG032207_0001 MWB027_WG032207_0002 7C27024-BLK1	J (all detects)	P
7C29027-BS1	2-Butanone 1,2-Dibromo-3-chloropropane 2-Hexanone 4-Methyl-2-pentanone	268 (40-140) 142 (50-135) 248 (45-140) 170 (45-140)	CMW002_WG032207_0001 7C29027-BLK1	J (all detects) J (all detects) J (all detects) J (all detects)	P

IX. Regional Quality Assurance and Quality Control

Not applicable.

X. Internal Standards

All internal standard areas and retention times were within QC limits.

XI. Target Compound Identifications

Raw data were not reviewed for this SDG.

XII. Compound Quantitation and CRQLs

Raw data were not reviewed for this SDG.

XIII. Tentatively Identified Compounds (TICs)

Raw data were not reviewed for this SDG.

XIV. System Performance

Raw data were not reviewed for this SDG.

XV. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

XVI. Field Duplicates

Samples MWB027_WG032207_0001 and MWB027_WG032207_0002 were identified as field duplicates. No volatiles were detected in any of the samples with the following exceptions:

Compound	Concentration (ug/L)		RPD
	MWB027_WG032207_0001	MWB027_WG032207_0002	
Chloroform	4.8	4.1	16
1,1-Dichloroethene	170	160	6
cis-1,2-Dichloroethene	120	110	9
trans-1,2-Dichloroethene	1.1	0.72	42
Tetrachloroethene	2.2	1.8	20
Toluene	1.2	0.91	27

Compound	Concentration (ug/L)		RPD
	MWB027_WG032207_0001	MWB027_WG032207_0002	
Trichlorofluoromethane	22	19	15
Trichloroethene	400	370	8
1,1-Dichloroethane	1.0U	0.3	200

XVII. Field Blanks

No field blanks were identified in this SDG.

Boeing Realty Corp., Bldg C-6 Torrance
Volatiles - Data Qualification Summary - SDG IQC2470

SDG	Sample	Compound	Flag	A or P	Reason
IQC2470	MWB028_WG032207_0001 MWB027_WG032207_0001 MWB027_WG032207_0002 CMW002_WG032207_0001	2-Butanone	J (all detects) UJ (all non-detects)	A	Initial calibration (RRF)
IQC2470	MWB028_WG032207_0001 MWB027_WG032207_0002 CMW002_WG032207_0001	Acetone	J (all detects) UJ (all non-detects)	A	Continuing calibration (%D)
IQC2470	CMW002_WG032207_0001	Tetrahydrofuran 2-Butanone 4-Methyl-2-pentanone 2-Hexanone	J (all detects) UJ (all non-detects)	A	Continuing calibration (%D)
IQC2470	MWB028_WG032207_0001 MWB027_WG032207_0001 MWB027_WG032207_0002	2-Butanone	J (all detects)	P	Laboratory control samples (%R)
IQC2470	MWB028_WG032207_0001 MWB027_WG032207_0002	Acetone	J (all detects)	P	Laboratory control samples (%R)
IQC2470	CMW002_WG032207_0001	2-Butanone 1,2-Dibromo-3-chloropropane 2-Hexanone 4-Methyl-2-pentanone	J (all detects) J (all detects) J (all detects) J (all detects)	P	Laboratory control samples (%R)

Boeing Realty Corp., Bldg C-6 Torrance
Volatiles - Laboratory Blank Data Qualification Summary - SDG IQC2470

SDG	Sample	Compound TIC (RT in minutes)	Modified Final Concentration	A or P
IQC2470	MWB028_WG032207_0001	Toluene	1.0U ug/L	A
IQC2470	MWB027_WG032207_0001	Toluene	1.2U ug/L	A
IQC2470	MWB027_WG032207_0002	Toluene	1.0U ug/L	A

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ANALYTICAL TESTING CORPORATION

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TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IQC2470

Sampled: 03/22/07
Received: 03/22/07

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC2470-05 (MWB028_WG032207_0001 - Water)									
Reporting Units: ug/l									
Acetone	EPA 8260B	7C27024	4.5	10	ND	4J 1	03/27/07	03/27/07	L
Benzene	EPA 8260B	7C27024	0.28	1.0	0.73	1	03/27/07	03/27/07	J
Bromobenzene	EPA 8260B	7C27024	0.27	1.0	ND	1	03/27/07	03/27/07	
Bromochloromethane	EPA 8260B	7C27024	0.32	1.0	ND	1	03/27/07	03/27/07	
Bromodichloromethane	EPA 8260B	7C27024	0.30	1.0	ND	1	03/27/07	03/27/07	
Bromoform	EPA 8260B	7C27024	0.40	1.0	ND	1	03/27/07	03/27/07	
Bromomethane	EPA 8260B	7C27024	0.42	1.0	ND	1	03/27/07	03/27/07	
2-Butanone (MEK)	EPA 8260B	7C27024	3.8	5.0	ND	4J 1	03/27/07	03/27/07	L
n-Butylbenzene	EPA 8260B	7C27024	0.37	1.0	ND	1	03/27/07	03/27/07	
sec-Butylbenzene	EPA 8260B	7C27024	0.25	1.0	ND	1	03/27/07	03/27/07	
tert-Butylbenzene	EPA 8260B	7C27024	0.22	1.0	ND	1	03/27/07	03/27/07	
Carbon Disulfide	EPA 8260B	7C27024	0.48	1.0	ND	1	03/27/07	03/27/07	
Carbon tetrachloride	EPA 8260B	7C27024	0.28	0.50	ND	1	03/27/07	03/27/07	
Chlorobenzene	EPA 8260B	7C27024	0.36	1.0	ND	1	03/27/07	03/27/07	
Chloroethane	EPA 8260B	7C27024	0.40	2.0	ND	1	03/27/07	03/27/07	
Chloroform	EPA 8260B	7C27024	0.33	1.0	3.0	1	03/27/07	03/27/07	
Chloromethane	EPA 8260B	7C27024	0.40	2.0	ND	1	03/27/07	03/27/07	
2-Chlorotoluene	EPA 8260B	7C27024	0.28	1.0	ND	1	03/27/07	03/27/07	
4-Chlorotoluene	EPA 8260B	7C27024	0.29	1.0	ND	1	03/27/07	03/27/07	
1,2-Dibromo-3-chloropropane	EPA 8260B	7C27024	0.97	2.0	ND	1	03/27/07	03/27/07	
Dibromochloromethane	EPA 8260B	7C27024	0.28	1.0	ND	1	03/27/07	03/27/07	
1,2-Dibromoethane (EDB)	EPA 8260B	7C27024	0.40	1.0	ND	1	03/27/07	03/27/07	
1,4-Dichlorobenzene	EPA 8260B	7C27024	0.37	1.0	ND	1	03/27/07	03/27/07	
1,2-Dichlorobenzene	EPA 8260B	7C27024	0.32	1.0	ND	1	03/27/07	03/27/07	
1,3-Dichlorobenzene	EPA 8260B	7C27024	0.35	1.0	ND	1	03/27/07	03/27/07	
Dichlorodifluoromethane	EPA 8260B	7C27024	0.79	1.0	ND	1	03/27/07	03/27/07	
1,2-Dichloroethane	EPA 8260B	7C27024	0.28	0.50	0.53	1	03/27/07	03/27/07	
1,1-Dichloroethane	EPA 8260B	7C27024	0.27	1.0	8.8	1	03/27/07	03/27/07	
1,1-Dichloroethene	EPA 8260B	7C27024	0.42	1.0	280	1	03/27/07	03/27/07	
cis-1,2-Dichloroethene	EPA 8260B	7C27024	0.32	1.0	17	1	03/27/07	03/27/07	
trans-1,2-Dichloroethene	EPA 8260B	7C27024	0.27	1.0	6.0	1	03/27/07	03/27/07	
1,2-Dichloropropane	EPA 8260B	7C27024	0.35	1.0	ND	1	03/27/07	03/27/07	
2,2-Dichloropropane	EPA 8260B	7C27024	0.34	1.0	ND	1	03/27/07	03/27/07	
cis-1,3-Dichloropropene	EPA 8260B	7C27024	0.22	0.50	ND	1	03/27/07	03/27/07	
1,1-Dichloropropene	EPA 8260B	7C27024	0.28	1.0	ND	1	03/27/07	03/27/07	
trans-1,3-Dichloropropene	EPA 8260B	7C27024	0.32	0.50	ND	1	03/27/07	03/27/07	
Ethylbenzene	EPA 8260B	7C27024	0.25	1.0	ND	1	03/27/07	03/27/07	
Hexachlorobutadiene	EPA 8260B	7C27024	0.38	1.0	ND	1	03/27/07	03/27/07	
2-Hexanone	EPA 8260B	7C27024	2.6	6.0	ND	1	03/27/07	03/27/07	
Iodomethane	EPA 8260B	7C27024	1.0	2.0	ND	1	03/27/07	03/27/07	
Isopropylbenzene	EPA 8260B	7C27024	0.25	1.0	ND	1	03/27/07	03/27/07	

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TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IQC2470

Sampled: 03/22/07
Received: 03/22/07

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC2470-05 (MWB028_WG032207_0001 - Water) - cont.									
Reporting Units: ug/l									
p-Isopropyltoluene	EPA 8260B	7C27024	0.28	1.0	ND	1	03/27/07	03/27/07	
Methyl-tert-butyl Ether (MTBE)	EPA 8260B	7C27024	0.32	1.0	ND	1	03/27/07	03/27/07	
Methylene chloride	EPA 8260B	7C27024	0.95	1.0	ND	1	03/27/07	03/27/07	
4-Methyl-2-pentanone (MIBK)	EPA 8260B	7C27024	3.5	5.0	ND	1	03/27/07	03/27/07	
n-Propylbenzene	EPA 8260B	7C27024	0.27	1.0	ND	1	03/27/07	03/27/07	
Styrene	EPA 8260B	7C27024	0.16	1.0	ND	1	03/27/07	03/27/07	
1,1,1,2-Tetrachloroethane	EPA 8260B	7C27024	0.27	1.0	ND	1	03/27/07	03/27/07	
1,1,2,2-Tetrachloroethane	EPA 8260B	7C27024	0.24	1.0	ND	1	03/27/07	03/27/07	
Tetrachloroethene	EPA 8260B	7C27024	0.32	1.0	1.9	1	03/27/07	03/27/07	
Tetrahydrofuran (THF)	EPA 8260B	7C27024	3.5	10	ND	1	03/27/07	03/27/07	
Toluene	EPA 8260B	7C27024	0.36	1.0	0.84	1, 0.4	03/27/07	03/27/07	B, J
1,2,3-Trichlorobenzene	EPA 8260B	7C27024	0.30	1.0	ND	1	03/27/07	03/27/07	
1,2,4-Trichlorobenzene	EPA 8260B	7C27024	0.48	1.0	ND	1	03/27/07	03/27/07	
1,1,2-Trichloroethane	EPA 8260B	7C27024	0.30	1.0	ND	1	03/27/07	03/27/07	
1,1,1-Trichloroethane	EPA 8260B	7C27024	0.30	1.0	ND	1	03/27/07	03/27/07	
Trichlorofluoromethane	EPA 8260B	7C27024	0.34	2.0	ND	1	03/27/07	03/27/07	
1,2,3-Trichloropropane	EPA 8260B	7C27024	0.40	1.0	ND	1	03/27/07	03/27/07	
1,2,4-Trimethylbenzene	EPA 8260B	7C27024	0.23	1.0	ND	1	03/27/07	03/27/07	
1,3,5-Trimethylbenzene	EPA 8260B	7C27024	0.26	1.0	ND	1	03/27/07	03/27/07	
Vinyl acetate	EPA 8260B	7C27024	1.7	6.0	ND	1	03/27/07	03/27/07	
Vinyl chloride	EPA 8260B	7C27024	0.30	0.50	ND	1	03/27/07	03/27/07	
Xylenes, Total	EPA 8260B	7C27024	0.90	1.0	ND	1	03/27/07	03/27/07	
Surrogate: 4-Bromofluorobenzene (80-120%)									102 %
Surrogate: Dibromofluoromethane (80-120%)									107 %
Surrogate: Toluene-d8 (80-120%)									102 %

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TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IQC2470

Sampled: 03/22/07
Received: 03/22/07

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC2470-05RE1 (MWB028_WG032207_0001 - Water) - cont.									
Reporting Units: ug/l									
Trichloroethene	EPA 8260B	7C28004	2.6	10	780	10	03/28/07	03/28/07	
Surrogate: 4-Bromofluorobenzene (80-120%)					106 %				
Surrogate: Dibromofluoromethane (80-120%)					107 %				
Surrogate: Toluene-d8 (80-120%)					110 %				

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MAR 28 2007

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TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IQC2470

Sampled: 03/22/07
Received: 03/22/07

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC2470-06 (MWB027_WG032207_0001 - Water)									
Reporting Units: ug/l									
Benzene	EPA 8260B	7C27024	0.28	1.0	ND	1	03/27/07	03/28/07	
Bromobenzene	EPA 8260B	7C27024	0.27	1.0	ND	1	03/27/07	03/28/07	
Bromochloromethane	EPA 8260B	7C27024	0.32	1.0	ND	1	03/27/07	03/28/07	
Bromodichloromethane	EPA 8260B	7C27024	0.30	1.0	ND	1	03/27/07	03/28/07	
Bromoform	EPA 8260B	7C27024	0.40	1.0	ND	1	03/27/07	03/28/07	
Bromomethane	EPA 8260B	7C27024	0.42	1.0	ND	1	03/27/07	03/28/07	
2-Butanone (MEK)	EPA 8260B	7C27024	3.8	5.0	ND	1	03/27/07	03/28/07	L
n-Butylbenzene	EPA 8260B	7C27024	0.37	1.0	ND	1	03/27/07	03/28/07	
sec-Butylbenzene	EPA 8260B	7C27024	0.25	1.0	ND	1	03/27/07	03/28/07	
tert-Butylbenzene	EPA 8260B	7C27024	0.22	1.0	ND	1	03/27/07	03/28/07	
Carbon Disulfide	EPA 8260B	7C27024	0.48	1.0	ND	1	03/27/07	03/28/07	
Carbon tetrachloride	EPA 8260B	7C27024	0.28	0.50	ND	1	03/27/07	03/28/07	
Chlorobenzene	EPA 8260B	7C27024	0.36	1.0	ND	1	03/27/07	03/28/07	
Chloroethane	EPA 8260B	7C27024	0.40	2.0	ND	1	03/27/07	03/28/07	
Chloroform	EPA 8260B	7C27024	0.33	1.0	4.8	1	03/27/07	03/28/07	
Chloromethane	EPA 8260B	7C27024	0.40	2.0	ND	1	03/27/07	03/28/07	
2-Chlorotoluene	EPA 8260B	7C27024	0.28	1.0	ND	1	03/27/07	03/28/07	
4-Chlorotoluene	EPA 8260B	7C27024	0.29	1.0	ND	1	03/27/07	03/28/07	
1,2-Dibromo-3-chloropropane	EPA 8260B	7C27024	0.97	2.0	ND	1	03/27/07	03/28/07	
Dibromochloromethane	EPA 8260B	7C27024	0.28	1.0	ND	1	03/27/07	03/28/07	
1,2-Dibromoethane (EDB)	EPA 8260B	7C27024	0.40	1.0	ND	1	03/27/07	03/28/07	
1,4-Dichlorobenzene	EPA 8260B	7C27024	0.37	1.0	ND	1	03/27/07	03/28/07	
1,2-Dichlorobenzene	EPA 8260B	7C27024	0.32	1.0	ND	1	03/27/07	03/28/07	
1,3-Dichlorobenzene	EPA 8260B	7C27024	0.35	1.0	ND	1	03/27/07	03/28/07	
Dichlorodifluoromethane	EPA 8260B	7C27024	0.79	1.0	ND	1	03/27/07	03/28/07	
1,2-Dichloroethane	EPA 8260B	7C27024	0.28	0.50	ND	1	03/27/07	03/28/07	
1,1-Dichloroethane	EPA 8260B	7C27024	0.27	1.0	ND	1	03/27/07	03/28/07	
1,1-Dichloroethene	EPA 8260B	7C27024	0.42	1.0	170	1	03/27/07	03/28/07	
cis-1,2-Dichloroethene	EPA 8260B	7C27024	0.32	1.0	120	1	03/27/07	03/28/07	
trans-1,2-Dichloroethene	EPA 8260B	7C27024	0.27	1.0	1.1	1	03/27/07	03/28/07	
1,2-Dichloropropane	EPA 8260B	7C27024	0.35	1.0	ND	1	03/27/07	03/28/07	
2,2-Dichloropropane	EPA 8260B	7C27024	0.34	1.0	ND	1	03/27/07	03/28/07	
cis-1,3-Dichloropropene	EPA 8260B	7C27024	0.22	0.50	ND	1	03/27/07	03/28/07	
1,1-Dichloropropene	EPA 8260B	7C27024	0.28	1.0	ND	1	03/27/07	03/28/07	
trans-1,3-Dichloropropene	EPA 8260B	7C27024	0.32	0.50	ND	1	03/27/07	03/28/07	
Ethylbenzene	EPA 8260B	7C27024	0.25	1.0	ND	1	03/27/07	03/28/07	
Hexachlorobutadiene	EPA 8260B	7C27024	0.38	1.0	ND	1	03/27/07	03/28/07	
2-Hexanone	EPA 8260B	7C27024	2.6	6.0	ND	1	03/27/07	03/28/07	
Iodomethane	EPA 8260B	7C27024	1.0	2.0	ND	1	03/27/07	03/28/07	
Isopropylbenzene	EPA 8260B	7C27024	0.25	1.0	ND	1	03/27/07	03/28/07	
p-Isopropyltoluene	EPA 8260B	7C27024	0.28	1.0	ND	1	03/27/07	03/28/07	

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TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IQC2470

Sampled: 03/22/07
Received: 03/22/07

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC2470-06 (MWB027_WG032207_0001 - Water) - cont.									
Reporting Units: ug/l									
Methyl-tert-butyl Ether (MTBE)	EPA 8260B	7C27024	0.32	1.0	ND	1	03/27/07	03/28/07	
Methylene chloride	EPA 8260B	7C27024	0.95	1.0	ND	1	03/27/07	03/28/07	
4-Methyl-2-pentanone (MIBK)	EPA 8260B	7C27024	3.5	5.0	ND	1	03/27/07	03/28/07	
n-Propylbenzene	EPA 8260B	7C27024	0.27	1.0	ND	1	03/27/07	03/28/07	
Styrene	EPA 8260B	7C27024	0.16	1.0	ND	1	03/27/07	03/28/07	
1,1,1,2-Tetrachloroethane	EPA 8260B	7C27024	0.27	1.0	ND	1	03/27/07	03/28/07	
1,1,2,2-Tetrachloroethane	EPA 8260B	7C27024	0.24	1.0	ND	1	03/27/07	03/28/07	
Tetrachloroethene	EPA 8260B	7C27024	0.32	1.0	2.2	1	03/27/07	03/28/07	
Tetrahydrofuran (THF)	EPA 8260B	7C27024	3.5	10	ND	1	03/27/07	03/28/07	
Toluene	EPA 8260B	7C27024	0.36	1.0	1.2 U	1	03/27/07	03/28/07	B
1,2,3-Trichlorobenzene	EPA 8260B	7C27024	0.30	1.0	ND	1	03/27/07	03/28/07	
1,2,4-Trichlorobenzene	EPA 8260B	7C27024	0.48	1.0	ND	1	03/27/07	03/28/07	
1,1,2-Trichloroethane	EPA 8260B	7C27024	0.30	1.0	ND	1	03/27/07	03/28/07	
1,1,1-Trichloroethane	EPA 8260B	7C27024	0.30	1.0	ND	1	03/27/07	03/28/07	
Trichlorofluoromethane	EPA 8260B	7C27024	0.34	2.0	22	1	03/27/07	03/28/07	
1,2,3-Trichloropropane	EPA 8260B	7C27024	0.40	1.0	ND	1	03/27/07	03/28/07	
1,2,4-Trimethylbenzene	EPA 8260B	7C27024	0.23	1.0	ND	1	03/27/07	03/28/07	
1,3,5-Trimethylbenzene	EPA 8260B	7C27024	0.26	1.0	ND	1	03/27/07	03/28/07	
Vinyl acetate	EPA 8260B	7C27024	1.7	6.0	ND	1	03/27/07	03/28/07	
Vinyl chloride	EPA 8260B	7C27024	0.30	0.50	ND	1	03/27/07	03/28/07	
Xylenes, Total	EPA 8260B	7C27024	0.90	1.0	ND	1	03/27/07	03/28/07	
Surrogate: 4-Bromofluorobenzene (80-120%)					101 %				
Surrogate: Dibromofluoromethane (80-120%)					120 %				
Surrogate: Toluene-d8 (80-120%)					103 %				

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Project Manager

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TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IQC2470

Sampled: 03/22/07
Received: 03/22/07

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC2470-06RE1 (MWB027_WG032207_0001 - Water) - cont.									
Reporting Units: ug/l									
Trichloroethene	EPA 8260B	7C28012	2.6	10	400	10	03/28/07	03/28/07	
Surrogate: 4-Bromofluorobenzene (80-120%)					106 %				
Surrogate: Dibromofluoromethane (80-120%)					119 %				
Surrogate: Toluene-d8 (80-120%)					106 %				

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TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IQC2470

Sampled: 03/22/07
Received: 03/22/07

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC2470-06RE2 (MWB027_WG032207_0001 - Water) - cont.									
Reporting Units: ug/l									
Acetone	EPA 8260B	7C29029	4.5	10	ND	1	03/29/07	03/30/07	
Surrogate: 4-Bromofluorobenzene (80-120%)					98 %				
Surrogate: Dibromofluoromethane (80-120%)					96 %				
Surrogate: Toluene-d8 (80-120%)					99 %				

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Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IQC2470

Sampled: 03/22/07
Received: 03/22/07

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC2470-07 (MWB027_WG032207_0002 - Water)									
Reporting Units: ug/l									
Acetone	EPA 8260B	7C27024	4.5	10	ND	1	03/27/07	03/28/07	L
Benzene	EPA 8260B	7C27024	0.28	1.0	ND	1	03/27/07	03/28/07	
Bromobenzene	EPA 8260B	7C27024	0.27	1.0	ND	1	03/27/07	03/28/07	
Bromochloromethane	EPA 8260B	7C27024	0.32	1.0	ND	1	03/27/07	03/28/07	
Bromodichloromethane	EPA 8260B	7C27024	0.30	1.0	ND	1	03/27/07	03/28/07	
Bromoform	EPA 8260B	7C27024	0.40	1.0	ND	1	03/27/07	03/28/07	
Bromomethane	EPA 8260B	7C27024	0.42	1.0	ND	1	03/27/07	03/28/07	
2-Butanone (MEK)	EPA 8260B	7C27024	3.8	5.0	ND	1	03/27/07	03/28/07	L
n-Butylbenzene	EPA 8260B	7C27024	0.37	1.0	ND	1	03/27/07	03/28/07	
sec-Butylbenzene	EPA 8260B	7C27024	0.25	1.0	ND	1	03/27/07	03/28/07	
tert-Butylbenzene	EPA 8260B	7C27024	0.22	1.0	ND	1	03/27/07	03/28/07	
Carbon Disulfide	EPA 8260B	7C27024	0.48	1.0	ND	1	03/27/07	03/28/07	
Carbon tetrachloride	EPA 8260B	7C27024	0.28	0.50	ND	1	03/27/07	03/28/07	
Chlorobenzene	EPA 8260B	7C27024	0.36	1.0	ND	1	03/27/07	03/28/07	
Chloroethane	EPA 8260B	7C27024	0.40	2.0	ND	1	03/27/07	03/28/07	
Chloroform	EPA 8260B	7C27024	0.33	1.0	4.1	1	03/27/07	03/28/07	
Chloromethane	EPA 8260B	7C27024	0.40	2.0	ND	1	03/27/07	03/28/07	
2-Chlorotoluene	EPA 8260B	7C27024	0.28	1.0	ND	1	03/27/07	03/28/07	
4-Chlorotoluene	EPA 8260B	7C27024	0.29	1.0	ND	1	03/27/07	03/28/07	
1,2-Dibromo-3-chloropropane	EPA 8260B	7C27024	0.97	2.0	ND	1	03/27/07	03/28/07	
Dibromochloromethane	EPA 8260B	7C27024	0.28	1.0	ND	1	03/27/07	03/28/07	
1,2-Dibromoethane (EDB)	EPA 8260B	7C27024	0.40	1.0	ND	1	03/27/07	03/28/07	
1,4-Dichlorobenzene	EPA 8260B	7C27024	0.37	1.0	ND	1	03/27/07	03/28/07	
1,2-Dichlorobenzene	EPA 8260B	7C27024	0.32	1.0	ND	1	03/27/07	03/28/07	
1,3-Dichlorobenzene	EPA 8260B	7C27024	0.35	1.0	ND	1	03/27/07	03/28/07	
Dichlorodifluoromethane	EPA 8260B	7C27024	0.79	1.0	ND	1	03/27/07	03/28/07	
1,2-Dichloroethane	EPA 8260B	7C27024	0.28	0.50	ND	1	03/27/07	03/28/07	
1,1-Dichloroethane	EPA 8260B	7C27024	0.27	1.0	0.30	1	03/27/07	03/28/07	J
1,1-Dichloroethene	EPA 8260B	7C27024	0.42	1.0	160	1	03/27/07	03/28/07	
cis-1,2-Dichloroethene	EPA 8260B	7C27024	0.32	1.0	110	1	03/27/07	03/28/07	
trans-1,2-Dichloroethene	EPA 8260B	7C27024	0.27	1.0	0.72	1	03/27/07	03/28/07	J
1,2-Dichloropropane	EPA 8260B	7C27024	0.35	1.0	ND	1	03/27/07	03/28/07	
2,2-Dichloropropane	EPA 8260B	7C27024	0.34	1.0	ND	1	03/27/07	03/28/07	
cis-1,3-Dichloropropene	EPA 8260B	7C27024	0.22	0.50	ND	1	03/27/07	03/28/07	
1,1-Dichloropropene	EPA 8260B	7C27024	0.28	1.0	ND	1	03/27/07	03/28/07	
trans-1,3-Dichloropropene	EPA 8260B	7C27024	0.32	0.50	ND	1	03/27/07	03/28/07	
Ethylbenzene	EPA 8260B	7C27024	0.25	1.0	ND	1	03/27/07	03/28/07	
Hexachlorobutadiene	EPA 8260B	7C27024	0.38	1.0	ND	1	03/27/07	03/28/07	
2-Hexanone	EPA 8260B	7C27024	2.6	6.0	ND	1	03/27/07	03/28/07	
Iodomethane	EPA 8260B	7C27024	1.0	2.0	ND	1	03/27/07	03/28/07	
Isopropylbenzene	EPA 8260B	7C27024	0.25	1.0	ND	1	03/27/07	03/28/07	

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TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IQC2470

Sampled: 03/22/07
Received: 03/22/07

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC2470-07 (MWB027_WG032207_0002 - Water) - cont.									
Reporting Units: ug/l									
p-Isopropyltoluene	EPA 8260B	7C27024	0.28	1.0	ND	1	03/27/07	03/28/07	
Methyl-tert-butyl Ether (MTBE)	EPA 8260B	7C27024	0.32	1.0	ND	1	03/27/07	03/28/07	
Methylene chloride	EPA 8260B	7C27024	0.95	1.0	ND	1	03/27/07	03/28/07	
4-Methyl-2-pentanone (MIBK)	EPA 8260B	7C27024	3.5	5.0	ND	1	03/27/07	03/28/07	
n-Propylbenzene	EPA 8260B	7C27024	0.27	1.0	ND	1	03/27/07	03/28/07	
Styrene	EPA 8260B	7C27024	0.16	1.0	ND	1	03/27/07	03/28/07	
1,1,1,2-Tetrachloroethane	EPA 8260B	7C27024	0.27	1.0	ND	1	03/27/07	03/28/07	
1,1,2,2-Tetrachloroethane	EPA 8260B	7C27024	0.24	1.0	ND	1	03/27/07	03/28/07	
Tetrachloroethene	EPA 8260B	7C27024	0.32	1.0	1.8	1	03/27/07	03/28/07	
Tetrahydrofuran (THF)	EPA 8260B	7C27024	3.5	10	ND	1	03/27/07	03/28/07	
Toluene	EPA 8260B	7C27024	0.36	1.0	0.91	1.04	03/27/07	03/28/07	B, J
1,2,3-Trichlorobenzene	EPA 8260B	7C27024	0.30	1.0	ND	1	03/27/07	03/28/07	
1,2,4-Trichlorobenzene	EPA 8260B	7C27024	0.48	1.0	ND	1	03/27/07	03/28/07	
1,1,2-Trichloroethane	EPA 8260B	7C27024	0.30	1.0	ND	1	03/27/07	03/28/07	
1,1,1-Trichloroethane	EPA 8260B	7C27024	0.30	1.0	ND	1	03/27/07	03/28/07	
Trichlorofluoromethane	EPA 8260B	7C27024	0.34	2.0	19	1	03/27/07	03/28/07	
1,2,3-Trichloropropane	EPA 8260B	7C27024	0.40	1.0	ND	1	03/27/07	03/28/07	
1,2,4-Trimethylbenzene	EPA 8260B	7C27024	0.23	1.0	ND	1	03/27/07	03/28/07	
1,3,5-Trimethylbenzene	EPA 8260B	7C27024	0.26	1.0	ND	1	03/27/07	03/28/07	
Vinyl acetate	EPA 8260B	7C27024	1.7	6.0	ND	1	03/27/07	03/28/07	
Vinyl chloride	EPA 8260B	7C27024	0.30	0.50	ND	1	03/27/07	03/28/07	
Xylenes, Total	EPA 8260B	7C27024	0.90	1.0	ND	1	03/27/07	03/28/07	
Surrogate: 4-Bromofluorobenzene (80-120%)									102 %
Surrogate: Dibromofluoromethane (80-120%)									119 %
Surrogate: Toluene-d8 (80-120%)									102 %

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TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IQC2470

Sampled: 03/22/07
Received: 03/22/07

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC2470-07RE1 (MWB027_WG032207_0002 - Water) - cont.									
Reporting Units: ug/l									
Trichloroethene	EPA 8260B	7C28004	2.6	10	370	10	03/28/07	03/28/07	
Surrogate: 4-Bromofluorobenzene (80-120%)					103 %				
Surrogate: Dibromofluoromethane (80-120%)					112 %				
Surrogate: Toluene-d8 (80-120%)					110 %				

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Project ID: Boeing C-6 Torrance
EM2727
Report Number: IQC2470

Sampled: 03/22/07
Received: 03/22/07

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC2470-13 (CMW002_WG032207_0001 - Water)									
Reporting Units: ug/l									
Benzene	EPA 8260B	7C29027	0.28	1.0	60	1	03/29/07	03/29/07	
Bromobenzene	EPA 8260B	7C29027	0.27	1.0	ND	1	03/29/07	03/29/07	
Bromochloromethane	EPA 8260B	7C29027	0.32	1.0	ND	1	03/29/07	03/29/07	
Bromodichloromethane	EPA 8260B	7C29027	0.30	1.0	ND	1	03/29/07	03/29/07	
Bromoform	EPA 8260B	7C29027	0.40	1.0	ND	1	03/29/07	03/29/07	
Bromomethane	EPA 8260B	7C29027	0.42	1.0	ND	1	03/29/07	03/29/07	
2-Butanone (MEK)	EPA 8260B	7C29027	3.8	5.0	ND	1	03/29/07	03/29/07	C, L
n-Butylbenzene	EPA 8260B	7C29027	0.37	1.0	ND	1	03/29/07	03/29/07	
sec-Butylbenzene	EPA 8260B	7C29027	0.25	1.0	ND	1	03/29/07	03/29/07	
tert-Butylbenzene	EPA 8260B	7C29027	0.22	1.0	ND	1	03/29/07	03/29/07	
Carbon Disulfide	EPA 8260B	7C29027	0.48	1.0	ND	1	03/29/07	03/29/07	
Carbon tetrachloride	EPA 8260B	7C29027	0.28	0.50	ND	1	03/29/07	03/29/07	
Chloroethane	EPA 8260B	7C29027	0.40	2.0	ND	1	03/29/07	03/29/07	
Chloroform	EPA 8260B	7C29027	0.33	1.0	1.1	1	03/29/07	03/29/07	
Chloromethane	EPA 8260B	7C29027	0.40	2.0	ND	1	03/29/07	03/29/07	
2-Chlorotoluene	EPA 8260B	7C29027	0.28	1.0	ND	1	03/29/07	03/29/07	
4-Chlorotoluene	EPA 8260B	7C29027	0.29	1.0	ND	1	03/29/07	03/29/07	
1,2-Dibromo-3-chloropropane	EPA 8260B	7C29027	0.97	2.0	ND	1	03/29/07	03/29/07	L
Dibromochloromethane	EPA 8260B	7C29027	0.28	1.0	ND	1	03/29/07	03/29/07	
1,2-Dibromoethane (EDB)	EPA 8260B	7C29027	0.40	1.0	ND	1	03/29/07	03/29/07	
1,4-Dichlorobenzene	EPA 8260B	7C29027	0.37	1.0	9.2	1	03/29/07	03/29/07	
1,2-Dichlorobenzene	EPA 8260B	7C29027	0.32	1.0	1.4	1	03/29/07	03/29/07	
1,3-Dichlorobenzene	EPA 8260B	7C29027	0.35	1.0	ND	1	03/29/07	03/29/07	
Dichlorodifluoromethane	EPA 8260B	7C29027	0.79	1.0	ND	1	03/29/07	03/29/07	
1,2-Dichloroethane	EPA 8260B	7C29027	0.28	0.50	ND	1	03/29/07	03/29/07	
1,1-Dichloroethane	EPA 8260B	7C29027	0.27	1.0	ND	1	03/29/07	03/29/07	
1,1-Dichloroethene	EPA 8260B	7C29027	0.42	1.0	1.2	1	03/29/07	03/29/07	
cis-1,2-Dichloroethene	EPA 8260B	7C29027	0.32	1.0	3.3	1	03/29/07	03/29/07	
trans-1,2-Dichloroethene	EPA 8260B	7C29027	0.27	1.0	ND	1	03/29/07	03/29/07	
1,2-Dichloropropane	EPA 8260B	7C29027	0.35	1.0	ND	1	03/29/07	03/29/07	
2,2-Dichloropropane	EPA 8260B	7C29027	0.34	1.0	ND	1	03/29/07	03/29/07	
cis-1,3-Dichloropropene	EPA 8260B	7C29027	0.22	0.50	ND	1	03/29/07	03/29/07	
1,1-Dichloropropene	EPA 8260B	7C29027	0.28	1.0	ND	1	03/29/07	03/29/07	
trans-1,3-Dichloropropene	EPA 8260B	7C29027	0.32	0.50	ND	1	03/29/07	03/29/07	
Ethylbenzene	EPA 8260B	7C29027	0.25	1.0	1.1	1	03/29/07	03/29/07	
Hexachlorobutadiene	EPA 8260B	7C29027	0.38	1.0	ND	1	03/29/07	03/29/07	
2-Hexanone	EPA 8260B	7C29027	2.6	6.0	ND	1	03/29/07	03/29/07	C, L
Iodomethane	EPA 8260B	7C29027	1.0	2.0	ND	1	03/29/07	03/29/07	
Isopropylbenzene	EPA 8260B	7C29027	0.25	1.0	ND	1	03/29/07	03/29/07	
p-Isopropyltoluene	EPA 8260B	7C29027	0.28	1.0	ND	1	03/29/07	03/29/07	
Methyl-tert-butyl Ether (MTBE)	EPA 8260B	7C29027	0.32	1.0	ND	1	03/29/07	03/29/07	

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Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IQC2470

Sampled: 03/22/07
Received: 03/22/07

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC2470-13 (CMW002_WG032207_0001 - Water) - cont.									
Reporting Units: ug/l									
4-Methyl-2-pentanone (MIBK)	EPA 8260B	7C29027	3.5	5.0	ND UJ	1	03/29/07	03/29/07	L
n-Propylbenzene	EPA 8260B	7C29027	0.27	1.0	ND	1	03/29/07	03/29/07	
Styrene	EPA 8260B	7C29027	0.16	1.0	ND	1	03/29/07	03/29/07	
1,1,1,2-Tetrachloroethane	EPA 8260B	7C29027	0.27	1.0	ND	1	03/29/07	03/29/07	
1,1,2,2-Tetrachloroethane	EPA 8260B	7C29027	0.24	1.0	ND	1	03/29/07	03/29/07	
Tetrachloroethene	EPA 8260B	7C29027	0.32	1.0	1.8	1	03/29/07	03/29/07	
Tetrahydrofuran (THF)	EPA 8260B	7C29027	3.5	10	ND UJ	1	03/29/07	03/29/07	C, L
Toluene	EPA 8260B	7C29027	0.36	1.0	ND	1	03/29/07	03/29/07	
1,2,3-Trichlorobenzene	EPA 8260B	7C29027	0.30	1.0	ND	1	03/29/07	03/29/07	
1,2,4-Trichlorobenzene	EPA 8260B	7C29027	0.48	1.0	ND	1	03/29/07	03/29/07	
1,1,2-Trichloroethane	EPA 8260B	7C29027	0.30	1.0	ND	1	03/29/07	03/29/07	
1,1,1-Trichloroethane	EPA 8260B	7C29027	0.30	1.0	ND	1	03/29/07	03/29/07	
Trichlorofluoromethane	EPA 8260B	7C29027	0.34	2.0	ND	1	03/29/07	03/29/07	
1,2,3-Trichloropropane	EPA 8260B	7C29027	0.40	1.0	ND	1	03/29/07	03/29/07	
1,2,4-Trimethylbenzene	EPA 8260B	7C29027	0.23	1.0	ND	1	03/29/07	03/29/07	
1,3,5-Trimethylbenzene	EPA 8260B	7C29027	0.26	1.0	ND	1	03/29/07	03/29/07	
Vinyl acetate	EPA 8260B	7C29027	1.7	6.0	ND	1	03/29/07	03/29/07	
Vinyl chloride	EPA 8260B	7C29027	0.30	0.50	ND	1	03/29/07	03/29/07	
Xylenes, Total	EPA 8260B	7C29027	0.90	1.0	2.4	1	03/29/07	03/29/07	
Surrogate: 4-Bromofluorobenzene (80-120%)					92 %				
Surrogate: Dibromofluoromethane (80-120%)					98 %				
Surrogate: Toluene-d8 (80-120%)					97 %				

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Project Manager

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Project ID: Boeing C-6 Torrance
EM2727
Report Number: IQC2470

Sampled: 03/22/07
Received: 03/22/07

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC2470-13RE1 (CMW002_WG032207_0001 - Water) - cont.									
Reporting Units: ug/l									
Chlorobenzene	EPA 8260B	7C30007	18	50	7400	50	03/30/07	03/30/07	
Surrogate: 4-Bromofluorobenzene (80-120%)					89 %				
Surrogate: Dibromofluoromethane (80-120%)					99 %				
Surrogate: Toluene-d8 (80-120%)					98 %				

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Project Manager

At 5/10/07

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Project ID: Boeing C-6 Torrance
EM2727
Report Number: IQC2470

Sampled: 03/22/07
Received: 03/22/07

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC2470-13RE2 (CMW002_WG032207_0001 - Water) - cont.									
Reporting Units: ug/l									
Acetone	EPA 8260B	7C30007	22	50	ND <i>UJ</i>	5	03/30/07	03/30/07	
Methylene chloride	EPA 8260B	7C30007	4.8	5.0	ND	5	03/30/07	03/30/07	
Trichloroethene	EPA 8260B	7C30007	1.3	5.0	340	5	03/30/07	03/30/07	
Surrogate: 4-Bromofluorobenzene (80-120%)					96 %				
Surrogate: Dibromofluoromethane (80-120%)					99 %				
Surrogate: Toluene-d8 (80-120%)					98 %				

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Nicholas Marz
Project Manager

M251007

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LDC #: 16739B1
 SDG #: IQC2470
 Laboratory: Test America

VALIDATION COMPLETENESS WORKSHEET
 Tier 2

Date: 5/10/07
 Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: GC/MS Volatiles (EPA SW 846 Method 8260B)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: 3/22/07
II.	GC/MS Instrument performance check	A	
III.	Initial calibration	SW	% RSD, r ² 10.990
IV.	Continuing calibration	SW	
V.	Blanks	SW	
VI.	Surrogate spikes	A	
VII.	Matrix spike/Matrix spike duplicates	SA	CMW001 - WG032207-0001 MS IP
VIII.	Laboratory control samples	SW	LC
IX.	Regional Quality Assurance and Quality Control	N	
X.	Internal standards	A	
XI.	Target compound identification	N	
XII.	Compound quantitation/CRQLs	N	
XIII.	Tentatively identified compounds (TICs)	N	
XIV.	System performance	N	
XV.	Overall assessment of data	A	
XVI.	Field duplicates	SW	D = 2 + 3
XVII.	Field blanks	N	

Note: A = Acceptable
 N = Not provided/applicable
 SW = See worksheet
 ND = No compounds detected
 R = Rinsate
 FB = Field blank
 D = Duplicate
 TB = Trip blank
 EB = Equipment blank

Validated Samples: water

5	1	2 = 5 MWB028_WG032207_0001	11	1	7027024 - BLK 1	21		31
6	2	3 = 5 4 = F MWB027_WG032207_0001	12	2	7028004 - BLK 1	22		32
7	3	2 = 5 MWB027_WG032207_0002	13	3	7028012 - BLK 1	23		33
13	4	6 = DD, F, E, S CMW002_WG032207_0001	14	4	7029029 - BLK 1	24		34
	5	MWB028_WG032207_0001MS	15	5	7029027 - BLK 1	25		35
	6	MWB028_WG032207_0001MSD	16	6	7030007 - BLK 1	26		36
	7		17			27		37
	8		18			28		38
	9		19			29		39
	10		20			30		40

TARGET COMPOUND WORKSHEET

METHOD: VOA (EPA SW 846 Method 8260B)

A. Chloromethane*	U. 1,1,2-Trichloroethane	OO. 2,2-Dichloropropane	III. n-Butylbenzene	CCCC. 1-Chlorohexane
B. Bromomethane	V. Benzene	PP. Bromochloromethane	JJJ. 1,2-Dichlorobenzene	DDDD. Isopropyl alcohol
C. Vinyl chloride**	W. trans-1,3-Dichloropropene	QQ. 1,1-Dichloropropene	KKK. 1,2,4-Trichlorobenzene	EEEE. Acetonitrile
D. Chloroethane	X. Bromoform*	RR. Dibromomethane	LLL. Hexachlorobutadiene	FFFF. Acrolein
E. Methylene chloride	Y. 4-Methyl-2-pentanone	SS. 1,3-Dichloropropane	MMM. Naphthalene	GGGG. Acrylonitrile
F. Acetone	Z. 2-Hexanone	TT. 1,2-Dibromoethane	NNN. 1,2,3-Trichlorobenzene	HHHH. 1,4-Dioxane
G. Carbon disulfide	AA. Tetrachloroethane	UU. 1,1,1,2-Tetrachloroethane	OOO. 1,3,5-Trichlorobenzene	IIII. Isobutyl alcohol
H. 1,1-Dichloroethene**	BB. 1,1,2-Tetrachloroethane*	VV. Isopropylbenzene	PPP. trans-1,2-Dichloroethene	JJJJ. Methacrylonitrile
I. 1,1-Dichloroethane*	CC. Toluene**	WW. Bromobenzene	QQQ. cis-1,2-Dichloroethene	KKKK. Propionitrile
J. 1,2-Dichloroethene, total	DD. Chlorobenzene*	XX. 1,2,3-Trichloropropane	RRR. m,p-Xylenes	LLLL. Ethyl ether
K. Chloroform**	EE. Ethylbenzene**	YY. n-Propylbenzene	SSS. o-Xylene	MMMM. Benzyl chloride
L. 1,2-Dichloroethane	FF. Styrene	ZZ. 2-Chlorotoluene	TTT. 1,1,2-Trichloro-1,2,2-trifluoroethane	NNNN.
M. 2-Butanone	GG. Xylenes, total	AAA. 1,3,5-Trimethylbenzene	UUU. 1,2-Dichlorotetrafluoroethane	OOOO.
N. 1,1,1-Trichloroethane	HH. Vinyl acetate	BBB. 4-Chlorotoluene	VVV. 4-Ethyltoluene	PPPP.
O. Carbon tetrachloride	II. 2-Chloroethylvinyl ether	CCC. tert-Butylbenzene	WWW. Ethanol	QQQQ.
P. Bromodichloromethane	JJ. Dichlorodifluoromethane	DDD. 1,2,4-Trimethylbenzene	XXX. Di-isopropyl ether	RRRR.
Q. 1,2-Dichloropropane**	KK. Trichlorofluoromethane	EEE. sec-Butylbenzene	YYY. tert-Butanol	SSSS.
R. cis-1,3-Dichloropropene	LL. Methyl-tert-butyl ether	FFF. 1,3-Dichlorobenzene	ZZZ. tert-Butyl alcohol	TTTT.
S. Trichloroethane	MM. 1,2-Dibromo-3-chloropropane	GGG. p-Isopropyltoluene	AAAA. Ethyl tert-butyl ether	UUUU.
T. Dibromochloromethane	NN. Methyl ethyl ketone	HHH. 1,4-Dichlorobenzene	BBBB. tert-Amyl methyl ether	VVVV.

* = System performance check compounds (SPCC) for RRF ; ** = Calibration check compounds (CCC) for %RSD.

LDC #: 16739B1
 SDG #: per count

VALIDATION FINDINGS WORKSHEET
Blanks

Page: 1 of 2
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: GC/MS VOA (EPA SW 846 Method 8260B)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

- Y N N/A Was a method blank associated with every sample in this SDG?
- Y N N/A Was a method blank analyzed at least once every 12 hours for each matrix and concentration?
- Y N N/A Was there contamination in the method blanks? If yes, please see the qualifications below.

Blank analysis date: 3/27/07

Conc. units: ug/l Associated Samples: 1-7 3

Compound	Blank ID	Sample Identification			
Tetrahydrofuran Methylene chloride	7027024-BLK1	1	2	3	
Acetone	804	-	-	-	
	0.440	0.84/1.0	1.2/1.0	0.9/1.0	
CRQL					

Blank analysis date: 3/29/07
 Conc. units: ug/l

Associated Samples: 4 (ND)

Compound	Blank ID	Sample Identification			
Tetrahydrofuran Methylene chloride	7029027-BLK1				
Acetone	5.88				
CRQL					

All results were qualified using the criteria stated below except those circled.

Note: Common contaminants such as Methylene chloride, Acetone, 2-Butanone, Carbon disulfide and TICs that were detected in samples within ten times the associated method blank concentration were qualified as not detected, "U". Other contaminants within five times the method blank concentration were also qualified as not detected, "U".

LDC #: 16739B1
 SDG #: pu cover

VALIDATION FINDINGS WORKSHEET
Blanks

Page: 7 of 2
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: GC/MS VOA (EPA SW 846 Method 8260B)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

- Y N N/A Was a method blank associated with every sample in this SDG?
- Y N N/A Was a method blank analyzed at least once every 12 hours for each matrix and concentration?
- Y N N/A Was there contamination in the method blanks? If yes, please see the qualifications below.

Blank analysis date: 2/30/07 Associated Samples: 4 (ND)
 Conc. units: ug/L

Compound	Blank ID	Sample Identification
	<u>1030007</u>	<u>BK1</u>
Methylene chloride	<u>1.64</u>	
Acetone		
CRQL		

Blank analysis date: _____
 Conc. units: _____

Associated Samples: _____

Compound	Blank ID	Sample Identification
Methylene chloride		
Acetone		
CRQL		

All results were qualified using the criteria stated below except those circled.

Note: Common contaminants such as Methylene chloride, Acetone, 2-Butanone, Carbon disulfide and TICs that were detected in samples within ten times the associated method blank concentration were qualified as not detected, "U". Other contaminants within five times the method blank concentration were also qualified as not detected, "U".

LDC #: 16739B1
 SDG #: per cover

VALIDATION FINDINGS WORKSHEET
Field Duplicates

Page: 1 of 1
 Reviewer: [Signature]
 2nd reviewer: [Signature]

METHOD: GCMS VOA (EPA SW 846 Method 8260B)

Y N N/A Were field duplicate pairs identified in this SDG?
Y N N/A Were target compounds detected in the field duplicate pairs?

Compound	Concentration (ug/L)		RPD
	2	3	
K	4.8	4.1	16
H	170	160	6
DDD	120	110	9
PPP	1.1	0.72	42
AA	2.2	1.8	20

Compound	Concentration ()		RPD
CC	1.2	0.91	27
KK	22	19	15
S	400	370	8
I	1.0u	0.3	200

Compound	Concentration ()		RPD

Compound	Concentration ()		RPD

LDC Report# 16739C1

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Boeing Realty Corp., Bldg C-6 Torrance

Collection Date: March 27, 2007

LDC Report Date: May 10, 2007

Matrix: Water

Parameters: Volatiles

Validation Level: Tier 3

Laboratory: TestAmerica

Sample Delivery Group (SDG): IQC2895

Sample Identification

MWB019_WG032707_0001

Introduction

This data review covers one water sample listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 8260B for Volatiles.

This review follows a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Organic Data Review (October 1999) as there are no current guidelines for the method stated above.

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

Blank results are summarized in Section V.

Field duplicates are summarized in Section XVI.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- N Presumptive evidence of presence of the constituent.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. GC/MS Instrument Performance Check

Instrument performance was checked at 12 hour intervals.

All ion abundance requirements were met.

III. Initial Calibration

Initial calibration was performed using required standard concentrations.

Percent relative standard deviations (%RSD) were less than or equal to 15.0% for each individual compound and less than or equal to 30.0% for calibration check compounds (CCCs).

In the case where %RSD was greater than 15.0%, the laboratory used a calibration curve to evaluate the compound. All coefficients of determination (r^2) were greater than or equal to 0.990 .

For the purposes of technical evaluation, all compounds were evaluated against the 30.0% (%RSD) National Functional Guideline criteria. Unless noted above, all compounds were within the validation criteria.

Average relative response factors (RRF) for all volatile target compounds and system performance check compounds (SPCCs) were within method and validation criteria with the following exceptions:

Date	Compound	RRF (Limits)	Associated Samples	Flag	A or P
2/28/07	2-Butanone	0.037 (≥ 0.05)	All samples in SDG IQC2895	J (all detects) UJ (all non-detects)	A

IV. Continuing Calibration

Continuing calibration was performed at the required frequencies.

Percent differences (%D) between the initial calibration RRF and the continuing calibration RRF were within the method criteria of less than or equal to 20.0% for calibration check compounds (CCCs).

For the purposes of technical evaluation, all compounds were evaluated against the 25.0% (%D) National Functional Guideline criteria. Unless noted above, all compounds were within the validation criteria with the following exceptions:

Date	Compound	%D	Associated Samples	Flag	A or P
4/3/07	Acetone 2-Butanone 2,2-Dichloropropane 2-Hexanone	109.3 70.3 29.1 63.9	All samples in SDG IQC2895	J (all detects) UJ (all non-detects)	A

All of the continuing calibration RRF values were within method and validation criteria.

V. Blanks

Method blanks were reviewed for each matrix as applicable. No volatile contaminants were found in the method blanks with the following exceptions:

Method Blank ID	Analysis Date	Compound TIC (RT in minutes)	Concentration	Associated Samples
7D03010-BLK1	4/3/07	Tetrahydrofuran Trichloroethene	8.04 ug/L 0.480 ug/L	All samples in SDG IQC2895

Sample concentrations were compared to concentrations detected in the method blanks. The sample concentrations were either not detected or were significantly greater (> 10X for common contaminants, >5X for other contaminants) than the concentrations found in the associated method blanks with the following exceptions:

Sample	Compound TIC (RT in minutes)	Reported Concentration	Modified Final Concentration
MWB019_WG032707_0001 (10X)	Tetrahydrofuran	64 ug/L	100U ug/L

VI. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VII. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were not within QC limits. Since there were no associated samples, no data were qualified.

VIII. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) were within QC limits with the following exceptions:

LCS ID	Compound	%R (Limits)	Associated Samples	Flag	A or P
7D03010-BS1	Acetone 2-Butanone 1,2-Dibromo-3-chloropropane 2-Hexanone 4-Methyl-2-pentanone 1,1,2,2-Tetrachloroethane 1,2,3-Trichloropropane	263 (30-140) 237 (40-140) 138 (50-135) 211 (45-140) 144 (45-140) 134 (55-130) 136 (60-130)	All samples in SDG IQC2895	J (all detects) J (all detects) J (all detects) J (all detects) J (all detects) J (all detects) J (all detects)	P

IX. Regional Quality Assurance and Quality Control

Not applicable.

X. Internal Standards

All internal standard areas and retention times were within QC limits.

XI. Target Compound Identifications

All target compound identifications were within validation criteria.

XII. Compound Quantitation and CRQLs

All compound quantitation and CRQLs were within validation criteria.

XIII. Tentatively Identified Compounds (TICs)

Tentatively identified compounds were not reported by the laboratory.

XIV. System Performance

The system performance was acceptable.

XV. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

XVI. Field Duplicates

No field duplicates were identified in this SDG.

XVII. Field Blanks

No field blanks were identified in this SDG.

Boeing Realty Corp., Bldg C-6 Torrance
Volatiles - Data Qualification Summary - SDG IQC2895

SDG	Sample	Compound	Flag	A or P	Reason
IQC2895	MWB019_WG032707_0001	2-Butanone	J (all detects) UJ (all non-detects)	A	Initial calibration (RRF)
IQC2895	MWB019_WG032707_0001	Acetone 2-Butanone 2,2-Dichloropropane 2-Hexanone	J (all detects) UJ (all non-detects)	A	Continuing calibration (%D)
IQC2895	MWB019_WG032707_0001	Acetone 2-Butanone 1,2-Dibromo-3-chloropropane 2-Hexanone 4-Methyl-2-pentanone 1,1,2,2-Tetrachloroethane 1,2,3-Trichloropropane	J (all detects) J (all detects) J (all detects) J (all detects) J (all detects) J (all detects) J (all detects)	P	Laboratory control samples (%R)

Boeing Realty Corp., Bldg C-6 Torrance
Volatiles - Laboratory Blank Data Qualification Summary - SDG IQC2895

SDG	Sample	Compound TIC (RT in minutes)	Modified Final Concentration	A or P
IQC2895	MWB019_WG032707_0001 (10X)	Tetrahydrofuran	100U ug/L	A

TestAmerica

ANALYTICAL TESTING CORPORATION

17461 Derian Avenue, Suite 100, Irvine, CA 92614 (949) 261-1022 Fax: (949) 260-3297

TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IQC2895

Sampled: 03/27/07
Received: 03/27/07

VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC2895-07 (MWB019_WG032707_0001 - Water)									
Reporting Units: ug/l									
Chloroform	EPA 8260B	7D02027	16	50	3600	50	04/02/07	04/03/07	
Surrogate: 4-Bromofluorobenzene (80-120%)					102 %				
Surrogate: Dibromofluoromethane (80-120%)					116 %				
Surrogate: Toluene-d8 (80-120%)					110 %				
Sample ID: IQC2895-07RE1 (MWB019_WG032707_0001 - Water)									
Reporting Units: ug/l									
Acetone	EPA 8260B	7D03010	45	100	ND	UJ 10	04/03/07	04/03/07	C, L
Benzene	EPA 8260B	7D03010	2.8	10	ND	10	04/03/07	04/03/07	
Bromobenzene	EPA 8260B	7D03010	2.7	10	ND	10	04/03/07	04/03/07	
Bromochloromethane	EPA 8260B	7D03010	3.2	10	ND	10	04/03/07	04/03/07	
Bromodichloromethane	EPA 8260B	7D03010	3.0	10	ND	10	04/03/07	04/03/07	
Bromoform	EPA 8260B	7D03010	4.0	10	ND	10	04/03/07	04/03/07	
Bromomethane	EPA 8260B	7D03010	4.2	10	ND	10	04/03/07	04/03/07	
2-Butanone (MEK)	EPA 8260B	7D03010	38	50	ND	UJ 10	04/03/07	04/03/07	C, L
n-Butylbenzene	EPA 8260B	7D03010	3.7	10	ND	10	04/03/07	04/03/07	
sec-Butylbenzene	EPA 8260B	7D03010	2.5	10	ND	10	04/03/07	04/03/07	
tert-Butylbenzene	EPA 8260B	7D03010	2.2	10	ND	10	04/03/07	04/03/07	
Carbon Disulfide	EPA 8260B	7D03010	4.8	10	ND	10	04/03/07	04/03/07	
Carbon tetrachloride	EPA 8260B	7D03010	2.8	5.0	10	10	04/03/07	04/03/07	
Chlorobenzene	EPA 8260B	7D03010	3.6	10	ND	10	04/03/07	04/03/07	
Chloroethane	EPA 8260B	7D03010	4.0	20	ND	10	04/03/07	04/03/07	
Chloromethane	EPA 8260B	7D03010	4.0	20	ND	10	04/03/07	04/03/07	
2-Chlorotoluene	EPA 8260B	7D03010	2.8	10	ND	10	04/03/07	04/03/07	
4-Chlorotoluene	EPA 8260B	7D03010	2.9	10	ND	10	04/03/07	04/03/07	
1,2-Dibromo-3-chloropropane	EPA 8260B	7D03010	9.7	20	ND	10	04/03/07	04/03/07	L
Dibromochloromethane	EPA 8260B	7D03010	2.8	10	ND	10	04/03/07	04/03/07	
1,2-Dibromoethane (EDB)	EPA 8260B	7D03010	4.0	10	ND	10	04/03/07	04/03/07	
1,4-Dichlorobenzene	EPA 8260B	7D03010	3.7	10	ND	10	04/03/07	04/03/07	
1,2-Dichlorobenzene	EPA 8260B	7D03010	3.2	10	ND	10	04/03/07	04/03/07	
1,3-Dichlorobenzene	EPA 8260B	7D03010	3.5	10	ND	10	04/03/07	04/03/07	
Dichlorodifluoromethane	EPA 8260B	7D03010	7.9	10	ND	10	04/03/07	04/03/07	
1,2-Dichloroethane	EPA 8260B	7D03010	2.8	5.0	ND	10	04/03/07	04/03/07	
1,1-Dichloroethane	EPA 8260B	7D03010	2.7	10	ND	10	04/03/07	04/03/07	
1,1-Dichloroethene	EPA 8260B	7D03010	4.2	10	4.3	10	04/03/07	04/03/07	J
cis-1,2-Dichloroethene	EPA 8260B	7D03010	3.2	10	ND	10	04/03/07	04/03/07	
trans-1,2-Dichloroethene	EPA 8260B	7D03010	2.7	10	ND	10	04/03/07	04/03/07	
1,2-Dichloropropane	EPA 8260B	7D03010	3.5	10	ND	10	04/03/07	04/03/07	
2,2-Dichloropropane	EPA 8260B	7D03010	3.4	10	ND	UJ 10	04/03/07	04/03/07	
cis-1,3-Dichloropropene	EPA 8260B	7D03010	2.2	5.0	ND	10	04/03/07	04/03/07	
1,1-Dichloropropene	EPA 8260B	7D03010	2.8	10	ND	10	04/03/07	04/03/07	

TestAmerica - Irvine, CA
Nicholas Marz
Project Manager

6251007

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IQC2895 <Page 19 of 76>

TAIT Environmental/Boeing 701 N. Parkcenter Drive Santa Ana, CA 92705 Attention: Mehmet Pehlivan	Project ID: Boeing C-6 Torrance EM2727 Report Number: IQC2895	Sampled: 03/27/07 Received: 03/27/07
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VOLATILE ORGANICS by GC/MS (EPA 5030B/8260B)

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC2895-07RE1 (MWB019_WG032707_0001 - Water) - cont.									
Reporting Units: ug/l									
trans-1,3-Dichloropropene	EPA 8260B	7D03010	3.2	5.0	ND	10	04/03/07	04/03/07	
Ethylbenzene	EPA 8260B	7D03010	2.5	10	ND	10	04/03/07	04/03/07	
Hexachlorobutadiene	EPA 8260B	7D03010	3.8	10	ND	10	04/03/07	04/03/07	
2-Hexanone	EPA 8260B	7D03010	26	60	ND 45	10	04/03/07	04/03/07	C, L
Iodomethane	EPA 8260B	7D03010	10	20	ND	10	04/03/07	04/03/07	
Isopropylbenzene	EPA 8260B	7D03010	2.5	10	ND	10	04/03/07	04/03/07	
p-Isopropyltoluene	EPA 8260B	7D03010	2.8	10	ND	10	04/03/07	04/03/07	
Methyl-tert-butyl Ether (MTBE)	EPA 8260B	7D03010	3.2	10	ND	10	04/03/07	04/03/07	
Methylene chloride	EPA 8260B	7D03010	9.5	10	ND	10	04/03/07	04/03/07	
4-Methyl-2-pentanone (MIBK)	EPA 8260B	7D03010	35	50	ND	10	04/03/07	04/03/07	L
n-Propylbenzene	EPA 8260B	7D03010	2.7	10	ND	10	04/03/07	04/03/07	
Styrene	EPA 8260B	7D03010	1.6	10	ND	10	04/03/07	04/03/07	
1,1,1,2-Tetrachloroethane	EPA 8260B	7D03010	2.7	10	ND	10	04/03/07	04/03/07	
1,1,2,2-Tetrachloroethane	EPA 8260B	7D03010	2.4	10	ND	10	04/03/07	04/03/07	L
Tetrachloroethene	EPA 8260B	7D03010	3.2	10	150	10	04/03/07	04/03/07	
Tetrahydrofuran (THF)	EPA 8260B	7D03010	35	100	64 100/10	10	04/03/07	04/03/07	B, J
Toluene	EPA 8260B	7D03010	3.6	10	ND	10	04/03/07	04/03/07	
1,2,3-Trichlorobenzene	EPA 8260B	7D03010	3.0	10	ND	10	04/03/07	04/03/07	
1,2,4-Trichlorobenzene	EPA 8260B	7D03010	4.8	10	ND	10	04/03/07	04/03/07	
1,1,2-Trichloroethane	EPA 8260B	7D03010	3.0	10	ND	10	04/03/07	04/03/07	
1,1,1-Trichloroethane	EPA 8260B	7D03010	3.0	10	ND	10	04/03/07	04/03/07	
Trichloroethene	EPA 8260B	7D03010	2.6	10	160	10	04/03/07	04/03/07	
Trichlorofluoromethane	EPA 8260B	7D03010	3.4	20	ND	10	04/03/07	04/03/07	
1,2,3-Trichloropropane	EPA 8260B	7D03010	4.0	10	ND	10	04/03/07	04/03/07	L
1,2,4-Trimethylbenzene	EPA 8260B	7D03010	2.3	10	ND	10	04/03/07	04/03/07	
1,3,5-Trimethylbenzene	EPA 8260B	7D03010	2.6	10	ND	10	04/03/07	04/03/07	
Vinyl acetate	EPA 8260B	7D03010	17	60	ND	10	04/03/07	04/03/07	
Vinyl chloride	EPA 8260B	7D03010	3.0	5.0	ND	10	04/03/07	04/03/07	
Xylenes, Total	EPA 8260B	7D03010	9.0	10	ND	10	04/03/07	04/03/07	
Surrogate: 4-Bromofluorobenzene (80-120%)					105 %				
Surrogate: Dibromofluoromethane (80-120%)					111 %				
Surrogate: Toluene-d8 (80-120%)					109 %				

TestAmerica - Irvine, CA
Nicholas Marz
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8251007

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LDC #: 16739C1
 SDG #: IQC2895
 Laboratory: Test America

VALIDATION COMPLETENESS WORKSHEET
 Tier 3

Date: 5/10/07
 Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: GC/MS Volatiles (EPA SW 846 Method 8260B)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: 3/27/07
II.	GC/MS Instrument performance check	A	
III.	Initial calibration	SW	% RSD, r ² 10.990
IV.	Continuing calibration	SW	
V.	Blanks	SW	
VI.	Surrogate spikes	A	
VII.	Matrix spike/Matrix spike duplicates	SW	TMW-06-WG032707-0001MS1D
VIII.	Laboratory control samples	SW	LES
IX.	Regional Quality Assurance and Quality Control	N	
X.	Internal standards	A	
XI.	Target compound identification	A	
XII.	Compound quantitation/CRQLs	A	
XIII.	Tentatively identified compounds (TICs)	N	not reported
XIV.	System performance	A	
XV.	Overall assessment of data	A	
XVI.	Field duplicates	N	
XVII.	Field blanks	N	

Note: A = Acceptable
 N = Not provided/applicable
 SW = See worksheet
 ND = No compounds detected
 R = Rinsate
 FB = Field blank
 D = Duplicate
 TB = Trip blank
 EB = Equipment blank

Validated Samples:

1	2 = BBK water MWB019_WG032707_0001	11	7D03010-BLK	21		31	
2		12	7D02027-BLK	22		32	
3		13		23		33	
4		14		24		34	
5		15		25		35	
6		16		26		36	
7		17		27		37	
8		18		28		38	
9		19		29		39	
10		20		30		40	

DC #: 1673901
 G #: I 5 C 2 8 9 5

VALIDATION FINDINGS CHECKLIST

Page: 1 of 2
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

Method: Volatiles (EPA SW 846 Method 8260B)

Validation Area	Yes	No	NA	Findings/Comments
All technical holding times were met.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Cooler temperature criteria was met.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the BFB performance results reviewed and found to be within the specified criteria?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all samples analyzed within the 12 hour clock criteria?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Did the laboratory perform a 5 point calibration prior to sample analysis?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all percent relative standard deviations (%RSD) and relative response factors (RRF) within method criteria for all CCCs and SPCCs?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a curve fit used for evaluation?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Did the initial calibration meet the curve fit acceptance criteria of > 0.990?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all percent relative standard deviations (%RSD) ≤ 30% and relative response factors (RRF) > 0.05?	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
Was a continuing calibration standard analyzed at least once every 12 hours for each instrument?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all percent differences (%D) and relative response factors (RRF) within method criteria for all CCCs and SPCCs?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all percent differences (%D) ≤ 25% and relative response factors (RRF) ≥ 0.05?	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
Was a method blank associated with every sample in this SDG?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a method blank analyzed at least once every 12 hours for each matrix and concentration?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was there contamination in the method blanks? If yes, please see the Blanks validation completeness worksheet.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all surrogate %R within QC limits?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
If the percent recovery (%R) for one or more surrogates was out of QC limits, was a reanalysis performed to confirm samples with %R outside of criteria?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were a matrix spike (MS) and matrix spike duplicate (MSD) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD. Soil / Water.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a MS/MSD analyzed every 20 samples of each matrix?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the QC limits?	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
Was an LCS analyzed for this SDG?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	

DC #: 16739C1
 SDG #: 16C2895

VALIDATION FINDINGS CHECKLIST

Page: 2 of 2
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

Validation Area	Yes	No	NA	Findings/Comments
Was an LCS analyzed per analytical batch?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the QC limits?	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
Were performance evaluation (PE) samples performed?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were the performance evaluation (PE) samples within the acceptance limits?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were internal standard area counts within -50% or +100% of the associated calibration standard?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were retention times within + 30 seconds of the associated calibration standard?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were relative retention times (RRT's) within + 0.06 RRT units of the standard?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Did compound spectra meet specified EPA "Functional Guidelines" criteria?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were chromatogram peaks verified and accounted for?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the correct internal standard (IS), quantitation ion and relative response factor (RRF) used to quantitate the compound?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were compound quantitation and CRQLs adjusted to reflect all sample dilutions and dry weight factors applicable to level IV validation?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the major ions (> 10 percent relative intensity) in the reference spectrum evaluated in sample spectrum?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were relative intensities of the major ions within ± 20% between the sample and the reference spectra?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Did the raw data indicate that the laboratory performed a library search for all required peaks in the chromatograms (samples and blanks)?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
System performance was found to be acceptable.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Overall assessment of data was found to be acceptable.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Field duplicate pairs were identified in this SDG.	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
Target compounds were detected in the field duplicates.	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Field blanks were identified in this SDG.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Target compounds were detected in the field blanks.	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	

TARGET COMPOUND WORKSHEET

METHOD: VOA (EPA SW 846 Method 8260B)

A. Chloromethane*	S. Trichloroethene	KK. Trichlorofluoromethane	CCC. tert-Butylbenzene	UUU. 1,2-Dichlorotetrafluoroethane
B. Bromomethane	T. Dibromochloromethane	LL. Methyl-tert-butyl ether	DDD. 1,2,4-Trimethylbenzene	VVV. 4-Ethyltoluene
C. Vinyl chloride**	U. 1,1,2-Trichloroethane	MM. 1,2-Dibromo-3-chloropropane	EEE. sec-Butylbenzene	WWW. Ethanol
D. Chloroethane	V. Benzene	NN. Methyl ethyl ketone	FFF. 1,3-Dichlorobenzene	XXX. Diisopropyl ether
E. Methylene chloride	W. trans-1,3-Dichloropropene	OO. 2,2-Dichloropropane	GGG. p-Isopropyltoluene	YYY. tert-Butanol
F. Acetone	X. Bromoform*	PP. Bromochloromethane	HHH. 1,4-Dichlorobenzene	ZZZ. tert-Butyl alcohol
G. Carbon disulfide	Y. 4-Methyl-2-pentanone	QQ. 1,1-Dichloropropane	III. n-Butylbenzene	AAA. Ethyl tert-butyl ether
H. 1,1-Dichloroethene**	Z. 2-Hexanone	RR. Dibromomethane	JJJ. 1,2-Dichlorobenzene	BBB. tert-Amyl methyl ether
I. 1,1-Dichloroethane*	AA. Tetrachloroethane	SS. 1,3-Dichloropropane	KKK. 1,2,4-Trichlorobenzene	CCCC. 1-Chlorohexane
J. 1,2-Dichloroethane, total	BB. 1,1,2,2-Tetrachloroethane*	TT. 1,2-Dibromomethane	LLL. Hexachlorobutadiene	DDDD. Isopropyl alcohol
K. Chloroform**	CC. Toluene**	UU. 1,1,1,2-Tetrachloroethane	MMM. Naphthalene	EEEE. Acetonitrile
L. 1,2-Dichloroethane	DD. Chlorobenzene*	VV. Isopropylbenzene	NNN. 1,2,3-Trichlorobenzene	FFFF. Acrolein
M. 2-Butanone	EE. Ethylbenzene**	WW. Bromobenzene	OOO. 1,3,5-Trichlorobenzene	GGGG. Acrylonitrile
N. 1,1,1-Trichloroethane	FF. Styrene	XX. 1,2,3-Trichloropropane	PPP. trans-1,2-Dichloroethane	HHHH. 1,4-Dioxane
O. Carbon tetrachloride	GG. Xylenes, total	YY. n-Propylbenzene	QQQ. cis-1,2-Dichloroethane	IIII. Isobutyl alcohol
P. Bromodichloromethane	HH. Vinyl acetate	ZZ. 2-Chlorotoluene	RRR. m,p-Xylenes	JJJJ. Methacrylonitrile
Q. 1,2-Dichloropropane**	II. 2-Chloroethylvinyl ether	AAA. 1,3,5-Trimethylbenzene	SSS. o-Xylene	KKKK. Propionitrile
R. cis-1,3-Dichloropropene	JJ. Dichlorodifluoromethane	BBB. 4-Chlorotoluene	TTT. 1,1,2-Trichloro-1,2,2-trifluoroethane	LLLL

* = System performance check compounds (SPCC) for RRF ; ** = Calibration check compounds (CCC) for %RSD.

LDC #: 16 3901
 SDG #: I 0802 895

VALIDATION FINDINGS WORKSHEET
Initial Calibration Calculation Verification

Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: GC/MS VOA (EPA SW 846 Method 8260B)

The Relative Response Factor (RRF), average RRF, and percent relative standard deviation (%RSD) were recalculated for the compounds identified below using the following calculations:

RRF = $(A_i/C_i)/(A_s/C_s)$
 average RRF = sum of the RRFs/number of standards
 %RSD = $100 * (S/X)$
 A_i = Area of compound,
 C_i = Concentration of compound,
 S = Standard deviation of the RRFs
 X = Mean of the RRFs
 A_s = Area of associated internal standard
 C_s = Concentration of internal standard

#	Standard ID	Calibration Date	Compound (Reference Internal Standard)	Reported		Recalculated		Reported		Recalculated	
				RRF (25 std)	RRF (25 std)	RRF (25 std)	Average RRF (Initial)	Average RRF (Initial)	%RSD	%RSD	
1	14A L	2/28/07	Methylene chloride (1st internal standard)	0.452	0.452	0.474	0.474	7.78	7.78	7.78	7.78
			Trichlorethene (2nd internal standard)	0.336	0.336	0.331	0.331	5.92	5.92	5.92	5.92
			Toluene (3rd internal standard)	1.365	1.365	1.314	1.314	10.24	10.24	10.24	10.24
2			Ethyl Benzene (1st internal standard)	1.639	1.639	1.623	1.623	4.92	4.92	4.92	4.92
			Trichlorethene (2nd internal standard)								
			Toluene (3rd internal standard)								
3			Methylene chloride (1st internal standard)								
			Trichlorethene (2nd internal standard)								
			Toluene (3rd internal standard)								
4			Methylene chloride (1st internal standard)								
			Trichlorethene (2nd internal standard)								
			Toluene (3rd internal standard)								

Comments: Refer to Initial Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

DC #: 16739C1
SDG #: F 002895

VALIDATION FINDINGS WORKSHEET
Continuing Calibration Results Verification

Page: 1 of 1
Reviewer: [Signature]
2nd Reviewer: [Signature]

METHOD: GC/MS VOA (EPA SW 846 Method 8260B)

The percent difference (%D) of the initial calibration average Relative Response Factors (RRFs) and the continuing calibration RRFs were recalculated for the compounds identified below using the following calculation:

% Difference = $100 * (\text{ave. RRF} - \text{RRF}) / \text{ave. RRF}$
RRF = $(A_s)(C_s) / (A_s)(C_s)$

Where: ave. RRF = Initial calibration average RRF
RRF = continuing calibration RRF
A_s = Area of compound,
C_s = Concentration of compound,
A_i = Area of associated internal standard
C_i = Concentration of internal standard

#	Standard ID	Calibration Date	Compound (Reference Internal Standard)	Average RRF (Initial)	Reported		Recalculated	
					RRF (CC)	%D	RRF (CC)	%D
1	6.46	4/2/07	CHLOROPYRIFOS Methylene chloride (1st internal standard)	0.756	0.872	15.3	0.872	15.3
			Trichloroethene (2nd internal standard)					
			Toluene (3rd internal standard)					
2	9.13	4/3/07	Methylene chloride (1st internal standard)	0.474	0.468	1.3	0.468	1.3
			Trichloroethene (2nd internal standard)	0.351	0.357	7.9	0.357	7.9
			1,2-DCE Toluene (3rd internal standard)	1.314	1.403	6.8	1.403	6.8
3			Methylene chloride (1st internal standard)	1.623	1.767	8.9	1.767	8.9
			Trichloroethene (2nd internal standard)					
			Toluene (3rd internal standard)					
4			Methylene chloride (1st internal standard)					
			Trichloroethene (2nd internal standard)					
			Toluene (3rd internal standard)					

Comments: Refer to Continuing Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

CONCL:15B

LDC #: 16739C1
 SDG #: IQC2895

VALIDATION FINDINGS WORKSHEET
Surrogate Results Verification

Page: 1 of 1
 Reviewer: [Signature]
 2nd reviewer: [Signature]

METHOD: GC/MS VOA (EPA SW 846 Method 8260B)

The percent recoveries (%R) of surrogates were recalculated for the compounds identified below using the following calculation:

% Recovery: $SF/SS * 100$

Where: SF = Surrogate Found
 SS = Surrogate Spiked

Sample ID: #1 10x

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Toluene-d8	25.0	27.17	109	109	0
Bromofluorobenzene	↓	25.73	103	103	↓
1,2-Dichloroethane-d4					
Dibromofluoromethane	↓	27.66	111	111	↓

Sample ID: _____

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Toluene-d8					
Bromofluorobenzene					
1,2-Dichloroethane-d4					
Dibromofluoromethane					

Sample ID: _____

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Toluene-d8					
Bromofluorobenzene					
1,2-Dichloroethane-d4					
Dibromofluoromethane					

Sample ID: _____

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Toluene-d8					
Bromofluorobenzene					
1,2-Dichloroethane-d4					
Dibromofluoromethane					

Sample ID: _____

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Toluene-d8					
Bromofluorobenzene					
1,2-Dichloroethane-d4					
Dibromofluoromethane					

LDC #: 1673901
 SDG #: I 602895

VALIDATION FINDINGS WORKSHEET
Matrix Spike/Matrix Spike Duplicates Results Verification

Page: 1 of 1
 Reviewer: FF
 2nd Reviewer: R

METHOD: GC/MS VOA (EPA SW 846 Method 8260B)

The percent recoveries (%R) and Relative Percent Difference (RPD) of the matrix spike and matrix spike duplicate were recalculated for the compounds identified below using the following calculation:

$$\% \text{ Recovery} = 100 * (SSC - SC) / SA$$

$$RPD = |MSC - MSDC| * 2 / (MSC + MSDC)$$

Where: SSC = Spiked sample concentration SC = Sample concentration
 SA = Spike added MSC = Matrix spike percent recovery MSDC = Matrix spike duplicate percent recovery

MS/MSD sample: INW-06 - W9032707 - 0001 MS 1D

Compound	Spike Added (ug/L)		Sample Concentration (ug/L)	Spiked Sample Concentration (ug/L)		Matrix Spike Percent Recovery		Matrix Spike Duplicate Percent Recovery		MS/MSD RPD	
	MS	MSD		MS	MSD	Reported	Recalc.	Reported	Recalc.	Reported	Recalculated
	1,1-Dichloroethene	25.0		25.0	13	36.7	36.4	95	95	94	94
Trichloroethene			140	158	144	72	72	16	16	9	9
Benzene			ND	26.4	26.0	106	106	104	104	2	2
Toluene			ND	26.5	26.2	106	106	105	105	1	1
Chlorobenzene			ND	27.4	27.2	110	110	109	109	1	1

Comments: Refer to Matrix Spike/Matrix Spike Duplicates findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 16739101
 SDG #: 1962895

VALIDATION FINDINGS WORKSHEET
Laboratory Control Sample Results Verification

Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: GC/MS VOA (EPA SW 846 Method 8260B)

The percent recoveries (%R) and Relative Percent Difference (RPD) of the laboratory control sample and laboratory control sample duplicate (if applicable) were recalculated for the compounds identified below using the following calculation:

% Recovery = $100 * SSC/SA$ Where: SSC = Spiked sample concentration
 SA = Spike added

RPD = $100 * |LCS - LCSD| / (LCS + LCSD)$

LCS = Laboratory control sample percent recovery

LCSD = Laboratory control sample duplicate percent recovery

LCS ID: 7D03010 - B > 1

Compound	Spike Added (ug/L)		Spiked Sample Concentration (ug/L)		LCS		LCSD		Percent Recovery		Percent Recovery		RPD	
	LCS	LCSD	LCS	LCSD	Reported	Recalc.	Reported	Recalc.	Reported	Recalc.	Reported	Recalc.	Reported	Recalc.
1,1-Dichloroethene	25.0	NA	24.2	NA	97	97	97	97						
Trichloroethene			24.8		99	99								
Benzene			25.3		101	101								
Toluene			25.3		101	101								
Chlorobenzene			26.2		105	105					NA			

Comments: Refer to Laboratory Control Sample findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

DC #: 1673901
 SDG #: IRL2895

VALIDATION FINDINGS WORKSHEET
 Sample Calculation Verification

Page: bf/
 Reviewer: R
 2nd reviewer: A

METHOD: GC/MS VOA (EPA SW 846 Method 8260B)

Y / N / N/A

Were all reported results recalculated and verified for all level IV samples?

Y / N / N/A

Were all recalculated results for detected target compounds agree within 10.0% of the reported results?

Concentration = $\frac{(A_s)(L)(DF)}{(A_i)(RRF)(V_s)(\%S)}$

- A_s = Area of the characteristic ion (EICP) for the compound to be measured
- A_i = Area of the characteristic ion (EICP) for the specific internal standard
- L = Amount of internal standard added in nanograms (ng)
- RRF = Relative response factor of the calibration standard.
- V_s = Volume or weight of sample pruged in milliliters (ml) or grams (g).
- Df = Dilution factor.
- %S = Percent solids, applicable to soils and solid matrices only.

Example:

Sample I.D. #1

Conc. = $\frac{(1624)(25)(10)}{(545335)(0.423)}$
 = 911.668
 10.5 ug/L

#	Sample ID	Compound	Reported Concentration ()	Calculated Concentration ()	Qualification

**Boeing Realty Corp., Bldg C-6 Torrance
Data Validation Reports
LDC# 16739**

Maganese

LDC

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Boeing Realty Corp., Bldg C-6 Torrance
Collection Date: March 14, 2007
LDC Report Date: May 10, 2007
Matrix: Water
Parameters: Manganese
Validation Level: Tier 1
Laboratory: TestAmerica

Sample Delivery Group (SDG): IQC1612

Sample Identification

MWB013_WG031407_0001
MWC021_WG031407_0001
MWB013_WG031407_0001MS
MWB013_WG031407_0001MSD

Introduction

This data review covers 4 water samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 6010B for Manganese.

This review follows a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Inorganic Data Review (October 2004) as there are no current guidelines for the methods stated above.

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

Blanks are summarized in Section III.

Field duplicates are summarized in Section XIII.

Samples indicated by a double asterisk on the front cover underwent a Tier 2 review. A Tier 1 review was performed on all of the other samples. Raw data were not evaluated for the samples reviewed by Tier 1 and 2 criteria since this review is based on QC data.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- N Presumptive evidence of presence of the constituent.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. Calibration

Calibration data were not reviewed for Tier 1.

III. Blanks

Method blanks were reviewed for each matrix as applicable. No contaminant concentrations were found in the preparation blanks.

IV. ICP Interference Check Sample (ICS) Analysis

ICP interference check sample analysis data were not reviewed for Tier 1.

V. Matrix Spike Analysis

Matrix spike (MS) and matrix spike duplicate (MSD) samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

VI. Duplicate Sample Analysis

Duplicate (DUP) sample analyses were reviewed for each matrix as applicable.

VII. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) were within QC limits.

VIII. Internal Standards

ICP-MS was not utilized in this SDG.

IX. Furnace Atomic Absorption QC

Graphite furnace atomic absorption was not utilized in this SDG.

X. ICP Serial Dilution

ICP serial dilution was not performed for this SDG.

XI. Sample Result Verification

Raw data were not reviewed for this SDG.

XII. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

XIII. Field Duplicates

No field duplicates were identified in this SDG.

XIV. Field Blanks

No field blanks were identified in this SDG.

**Boeing Realty Corp., Bldg C-6 Torrance
Manganese - Data Qualification Summary - SDG IQC1612**

No Sample Data Qualified in this SDG

**Boeing Realty Corp., Bldg C-6 Torrance
Manganese - Laboratory Blank Data Qualification Summary - SDG IQC1612**

No Sample Data Qualified in this SDG

TestAmerica

ANALYTICAL TESTING CORPORATION

17461 Derian Avenue, Suite 100, Irvine, CA 92614 (949) 261-1022 Fax: (949) 260-3297

TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727-01
Report Number: IQC1612

Sampled: 03/14/07
Received: 03/14/07

DISSOLVED METALS

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC1612-05 (MWB013_WG031407_0001 - Water)									
Reporting Units: mg/l									
Manganese	EPA 6010B-Diss	7C14164	N/A	0.020	ND	1	03/14/07	03/15/07	
Sample ID: IQC1612-10 (MWC021_WG031407_0001 - Water)									
Reporting Units: mg/l									
Manganese	EPA 6010B-Diss	7C14164	N/A	0.020	0.030	1	03/14/07	03/15/07	
Sample ID: IQC1612-11 (TMW_10_WG031407_0001 - Water)									
Reporting Units: mg/l									
Manganese	EPA 6010B-Diss	7C14164	N/A	0.020	ND	1	03/14/07	03/15/07	
Sample ID: IQC1612-12 (TMW_15_WG031407_0001 - Water)									
Reporting Units: mg/l									
Manganese	EPA 6010B-Diss	7C14164	N/A	0.020	ND	1	03/14/07	03/15/07	

2651007

TestAmerica - Irvine, CA
Nicholas Marz
Project Manager

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LDC #: 16739A4
 SDG #: IQC1612
 Laboratory: Test America

VALIDATION COMPLETENESS WORKSHEET
 EPA Region 1 - Tier 1

Date: 5/10/07
 Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: Manganese (EPA SW 846 Method 6010B)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: 3/14/07
II.	Calibration	N	
III.	Blanks	A	PB
IV.	ICP Interference Check Sample (ICS) Analysis	N	
V.	Matrix Spike Analysis	A	3 us / usd
VI.	Duplicate Sample Analysis	N	
VII.	Laboratory Control Samples (LCS)	A	LCS
VIII.	Internal Standard (ICP-MS)	N	36.7 utilized
IX.	Furnace Atomic Absorption QC	N	
X.	ICP Serial Dilution	N	kit performed
XI.	Sample Result Verification	N	
XII.	Overall Assessment of Data	A	
XIII.	Field Duplicates	N	
XIV.	Field Blanks	N	

Note: A = Acceptable
 N = Not provided/applicable
 SW = See worksheet
 ND = No compounds detected
 R = Rinsate
 FB = Field blank
 D = Duplicate
 TB = Trip blank
 EB = Equipment blank

Validated Samples: A2

1	MWB013_WG031407_0001	11		21		31	
2	MWC021_WG031407_0001	12		22		32	
3	MWB013_WG031407_0001MS	13		23		33	
4	MWB013_WG031407_0001MSD	14		24		34	
5	PB	15		25		35	
6		16		26		36	
7		17		27		37	
8		18		28		38	
9		19		29		39	
10		20		30		40	

Notes: _____

LDC Report# 16739B4

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Boeing Realty Corp., Bldg C-6 Torrance
Collection Date: March 22, 2007
LDC Report Date: May 10, 2007
Matrix: Water
Parameters: Manganese
Validation Level: Tier 2
Laboratory: TestAmerica

Sample Delivery Group (SDG): IQC2470

Sample Identification

CMW002_WG032207_0001

Introduction

This data review covers one water sample listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 6010B for Manganese.

This review follows a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Inorganic Data Review (October 2004) as there are no current guidelines for the methods stated above.

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

Blanks are summarized in Section III.

Field duplicates are summarized in Section XIII.

Raw data were not reviewed for this SDG. The review was based on QC data.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- N Presumptive evidence of presence of the constituent.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. Calibration

An initial calibration was performed.

The frequency and analysis criteria of the initial calibration verification (ICV) and continuing calibration verification (CCV) were met.

III. Blanks

Method blanks were reviewed for each matrix as applicable. No contaminant concentrations were found in the initial, continuing and preparation blanks.

IV. ICP Interference Check Sample (ICS) Analysis

The frequency of analysis was met.

The criteria for analysis were met.

V. Matrix Spike Analysis

Matrix spike (MS) and matrix spike duplicate (MSD) samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

VI. Duplicate Sample Analysis

Duplicate (DUP) sample analyses were reviewed for each matrix as applicable.

VII. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) were within QC limits.

VIII. Internal Standards

ICP-MS was not utilized in this SDG.

IX. Furnace Atomic Absorption QC

Graphite furnace atomic absorption was not utilized in this SDG.

X. ICP Serial Dilution

ICP serial dilution was not performed for this SDG.

XI. Sample Result Verification

Raw data were not reviewed for this SDG.

XII. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

XIII. Field Duplicates

No field duplicates were identified in this SDG.

XIV. Field Blanks

No field blanks were identified in this SDG.

**Boeing Realty Corp., Bldg C-6 Torrance
Manganese - Data Qualification Summary - SDG IQC2470**

No Sample Data Qualified in this SDG

**Boeing Realty Corp., Bldg C-6 Torrance
Manganese - Laboratory Blank Data Qualification Summary - SDG IQC2470**

No Sample Data Qualified in this SDG

TestAmerica

ANALYTICAL TESTING CORPORATION

17461 Derian Avenue, Suite 100, Irvine, CA 92614 (949) 261-1022 Fax: (949) 260-3297

TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IQC2470

Sampled: 03/22/07
Received: 03/22/07

DISSOLVED METALS

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC2470-08 (MWB003_WG032207_0001 - Water)									
Reporting Units: mg/l									
Manganese	EPA 6010B-Diss	7C23136	0.0070	0.020	0.31	1	03/23/07	03/24/07	
Sample ID: IQC2470-10 (WCC06S_WG032207_0001 - Water)									
Reporting Units: mg/l									
Manganese	EPA 6010B-Diss	7C23136	0.0070	0.020	4.5	1	03/23/07	03/24/07	
Sample ID: IQC2470-13 (CMW002_WG032207_0001 - Water)									
Reporting Units: mg/l									
Manganese	EPA 6010B-Diss	7C23136	0.0070	0.020	0.15	1	03/23/07	03/24/07	
Sample ID: IQC2470-14 (HRZCMW002_WG032207_0001 - Water)									
Reporting Units: mg/l									
Manganese	EPA 6010B-Diss	7C23136	0.0070	0.020	2.8	1	03/23/07	03/24/07	

TestAmerica - Irvine, CA
Nicholas Marz
Project Manager

Handwritten: 12/5/07

The results pertain only to the samples tested in the laboratory. This report shall not be reproduced, except in full, without written permission from TestAmerica.

IQC2470 <Page 51 of 98>

LDC #: 16739B4
 SDG #: IQC2470
 Laboratory: Test America

VALIDATION COMPLETENESS WORKSHEET
 EPA Region 1 - Tier 2

Date: 5/10/07
 Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: Manganese (EPA SW 846 Method 6010B)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: 3/22/07
II.	Calibration	A	
III.	Blanks	A	
IV.	ICP Interference Check Sample (ICS) Analysis	A	
V.	Matrix Spike Analysis	A	was done.
VI.	Duplicate Sample Analysis	N	
VII.	Laboratory Control Samples (LCS)	A	was
VIII.	Internal Standard (ICP-MS)	N	not utilized
IX.	Furnace Atomic Absorption QC	N	
X.	ICP Serial Dilution	N	not performed
XI.	Sample Result Verification	N	
XII.	Overall Assessment of Data	A	
XIII.	Field Duplicates	N	
XIV.	Field Blanks	N	

Note: A = Acceptable ND = No compounds detected D = Duplicate
 N = Not provided/applicable R = Rinstate TB = Trip blank
 SW = See worksheet FB = Field blank EB = Equipment blank

Validated Samples: [Signature]

1	CMW002_WG032207_0001	11		21		31	
2	PB	12		22		32	
3		13		23		33	
4		14		24		34	
5		15		25		35	
6		16		26		36	
7		17		27		37	
8		18		28		38	
9		19		29		39	
10		20		30		40	

Notes: _____

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Boeing Realty Corp., Bldg C-6 Torrance
Collection Date: March 27, 2007
LDC Report Date: May 10, 2007
Matrix: Water
Parameters: Manganese
Validation Level: Tier 3
Laboratory: TestAmerica
Sample Delivery Group (SDG): IQC2895

Sample Identification

MWB019_WG032707_0001
MWB019_WG032707_0001MS
MWB019_WG032707_0001MSD

Introduction

This data review covers 3 water samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 6010B for Manganese.

This review follows a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Inorganic Data Review (October 2004) as there are no current guidelines for the methods stated above.

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

Blanks are summarized in Section III.

Field duplicates are summarized in Section XIII.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- N Presumptive evidence of presence of the constituent.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. Calibration

An initial calibration was performed.

The frequency and analysis criteria of the initial calibration verification (ICV) and continuing calibration verification (CCV) were met.

III. Blanks

Method blanks were reviewed for each matrix as applicable. No contaminant concentrations were found in the initial, continuing and preparation blanks.

IV. ICP Interference Check Sample (ICS) Analysis

The frequency of analysis was met.

The criteria for analysis were met.

V. Matrix Spike Analysis

Matrix spike (MS) and matrix spike duplicate (MSD) samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

VI. Duplicate Sample Analysis

Duplicate (DUP) sample analyses were reviewed for each matrix as applicable.

VII. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) were within QC limits.

VIII. Internal Standard (ICP-MS)

ICP-MS was not utilized in this SDG.

IX. Furnace Atomic Absorption QC

Graphite furnace atomic absorption was not utilized in this SDG.

X. ICP Serial Dilution

ICP serial dilution was not performed for this SDG.

XI. Sample Result Verification

All sample result verifications were acceptable.

XII. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

XIII. Field Duplicates

No field duplicates were identified in this SDG.

XIV. Field Blanks

No field blanks were identified in this SDG.

**Boeing Realty Corp., Bldg C-6 Torrance
Manganese - Data Qualification Summary - SDG IQC2895**

No Sample Data Qualified in this SDG

**Boeing Realty Corp., Bldg C-6 Torrance
Manganese - Laboratory Blank Data Qualification Summary - SDG IQC2895**

No Sample Data Qualified in this SDG

TestAmerica

ANALYTICAL TESTING CORPORATION

17461 Derian Avenue, Suite 100, Irvine, CA 92614 (949) 261-1022 Fax: (949) 260-3297

TAIT Environmental/Boeing 701 N. Parkcenter Drive Santa Ana, CA 92705 Attention: Mehmet Pehlivan	Project ID: Boeing C-6 Torrance EM2727 Report Number: IQC2895	Sampled: 03/27/07 Received: 03/27/07
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DISSOLVED METALS

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC2895-07 (MWB019_WG032707_0001 - Water)									
Reporting Units: mg/l									
Manganese	EPA 6010B-Diss	7C28102	0.0070	0.020	ND	1	03/28/07	03/29/07	
Sample ID: IQC2895-08 (MWC017_WG032707_0001 - Water)									
Reporting Units: mg/l									
Manganese	EPA 6010B-Diss	7C28102	0.0070	0.020	0.082	1	03/28/07	03/29/07	
Sample ID: IQC2895-09 (IRZMW005_WG032707_0001 - Water)									
Reporting Units: mg/l									
Manganese	EPA 6010B-Diss	7C28102	0.0070	0.020	1.6	1	03/28/07	03/29/07	

TestAmerica - Irvine, CA
Nicholas Marz
Project Manager

Handwritten: 1/11/07

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IQC2895 <Page 25 of 76>

LDC #: 16739C4
 SDG #: IQC2895
 Laboratory: Test America

VALIDATION COMPLETENESS WORKSHEET
 EPA Region 1 - Tier 3

Date: 5/10/07
 Page: (of 1)
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: Manganese (EPA SW 846 Method 6010B)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: 3/29/07
II.	Calibration	A	
III.	Blanks	A	
IV.	ICP Interference Check Sample (ICS) Analysis	A	
V.	Matrix Spike Analysis	A	3 MS/MSD
VI.	Duplicate Sample Analysis	N	
VII.	Laboratory Control Samples (LCS)	A	Leg
VIII.	Internal Standard (ICP-MS)	N	3 not in the list
IX.	Furnace Atomic Absorption QC	N	
X.	ICP Serial Dilution	N	not performed
XI.	Sample Result Verification	A	
XII.	Overall Assessment of Data	A	
XIII.	Field Duplicates	N	
XIV.	Field Blanks	N	

Note: A = Acceptable
 N = Not provided/applicable
 SW = See worksheet
 ND = No compounds detected
 R = Rinsate
 FB = Field blank
 D = Duplicate
 TB = Trip blank
 EB = Equipment blank

Validated Samples: *h2*

1	MWB019_WG032707_0001	11		21		31	
2	MWB019_WG032707_0001MS	12		22		32	
3	MWB019_WG032707_0001MSD	13		23		33	
4	PB	14		24		34	
5		15		25		35	
6		16		26		36	
7		17		27		37	
8		18		28		38	
9		19		29		39	
10		20		30		40	

Notes: _____

LDC #: 673904
 SDG #: TA c-2895

VALIDATION FINDINGS CHECKLIST

Page: 1 of 2
 Reviewer: HY
 2nd Reviewer: ✓

Method: Metals (EPA SW 846 Method 6010/7000/6020)

Validation Area	Yes	No	NA	Findings/Comments
III: Instrumentation				
All technical holding times were met.	✓			
Cooler temperature criteria was met.	✓			
IV: Calibration				
Were all instruments calibrated daily, each set-up time?	✓			
Were the proper number of standards used?	✓			
Were all initial and continuing calibration verification %Rs within the 90-110% (80-120% for mercury and 85-115% for cyanide) QC limits?	✓			
Were all initial calibration correlation coefficients > 0.995? (Level IV only)	✓			
V: Quality Control				
Was a method blank associated with every sample in this SDG?	✓			
Was there contamination in the method blanks? If yes, please see the Blanks validation completeness worksheet.		✓		
VI: ICP Interference Check Samples				
Were ICP interference check samples performed daily?	✓			
Were the AB solution percent recoveries (%R) with the 80-120% QC limits?	✓			
VII: Matrix Spike/Matrix Spike Duplicate				
Were a matrix spike (MS) and duplicate (DUP) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD or MS/DUP. Soil / Water.	✓			
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the 75-125 QC limits? If the sample concentration exceeded the spike concentration by a factor of 4 or more, no action was taken.	✓			
Were the MS/MSD or duplicate relative percent differences (RPD) < 20% for waters and < 35% for soil samples? A control limit of +/- RL (+/- 2X RL for soil) was used for samples that were < 5X the RL, including when only one of the duplicate sample values were < 5X the RL.	✓			
VIII: Laboratory control samples				
Was an LCS analyzed for this SDG?	✓			
Was an LCS analyzed per extraction batch?	✓			
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the 80-120% QC limits for water samples and laboratory established QC limits for soils?	✓			
IX: Furnace Atomic Absorption QC				
If MSA was performed, was the correlation coefficients > 0.995?			✓	
Do all applicable analyses have duplicate injections? (Level IV only)			✓	
For sample concentrations > RL, are applicable duplicate injection RSD values < 20%? (Level IV only)			✓	
Were analytical spike recoveries within the 85-115% QC limits?			✓	

LDC #: 16739
 SDG #: 202895

VALIDATION FINDINGS CHECKLIST

Page: 2 of 2
 Reviewer: JM
 2nd Reviewer: AK

Validation Area	Yes	No	NA	Findings/Comments
Was an ICP serial dilution analyzed if analyte concentrations were > 50X the IDL?			✓	
Were all percent differences (%Ds) < 10%?			✓	
Was there evidence of negative interference? If yes, professional judgement will be used to qualify the data.			✓	
Were all the percent recoveries (%R) within the 30-120% of the intensity of the internal standard in the associated initial calibration?			✓	
If the %Rs were outside the criteria, was a reanalysis performed?			✓	
Were performance evaluation (PE) samples performed?			✓	
Were the performance evaluation (PE) samples within the acceptance limits?			✓	
Were RLs adjusted to reflect all sample dilutions and dry weight factors applicable to level IV validation?	✓			
Overall assessment of data was found to be acceptable.	✓			
Field duplicate pairs were identified in this SDG.		✓		
Target analytes were detected in the field duplicates.			✓	
Field blanks were identified in this SDG.		✓		
Target analytes were detected in the field blanks.			✓	

LDC #: 16739 04
 SDG #: 1022895

VALIDATION FINDINGS WORKSHEET
Initial and Continuing Calibration Calculation Verification

Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: Trace Metals (EPA SW 846 Method 6010/7000)

An initial and continuing calibration verification percent recovery (%R) was recalculated for each type of analysis using the following formula:

$$\%R = \frac{\text{Found}}{\text{True}} \times 100$$

Where: Found = concentration (in ug/L) of each analyte measured in the analysis of the ICV or CCV solution
 True = concentration (in ug/L) of each analyte in the ICV or CCV source

Standard ID	Type of Analysis	Element	Found (ug/L)	True (ug/L)	Recalculated		Reported		Acceptable (Y/N)
					%R		%R		
ICV	ICP (Initial calibration)	Mn	1.97	2	96		NR		Y
	GFAA (Initial calibration)								
	CVAA (Initial calibration)								
CCV	ICP (Continuing calibration)	Mn	1.03	1	103		NR		Y
	GFAA (Continuing calibration)								
	CVAA (Continuing calibration)								
	Cyanide (Initial calibration)								
	Cyanide (Continuing calibration)								

Comments: Refer to Calibration Verification findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

CALCLC 4SW

LDC #: 1673904
 SDG #: 1673904

Page: 1 of 1
 Reviewer: JK
 2nd Reviewer: JK

VALIDATION FINDINGS WORKSHEET
Level IV Recalculation Worksheet

METHOD: Trace Metals (EPA SW 846 Method 6010/7000)

Percent recoveries (%R) for an ICP interference check sample, a laboratory control sample and a matrix spike sample were recalculated using the following formula:

$$\%R = \frac{\text{Found}}{\text{True}} \times 100$$
 Where, Found = Concentration of each analyte measured in the analysis of the sample. For the matrix spike calculation,
 Found = SSR (spiked sample result) - SR (sample result).
 True = Concentration of each analyte in the source.

A sample and duplicate relative percent difference (RPD) was recalculated using the following formula:

$$RPD = \frac{|S-D|}{(S+D)/2} \times 100$$
 Where, S = Original sample concentration
 D = Duplicate sample concentration

An ICP serial dilution percent difference (%D) was recalculated using the following formula:

$$\%D = \frac{|I-SDR|}{I} \times 100$$
 Where, I = Initial Sample Result (mg/L)
 SDR = Serial Dilution Result (mg/L) (Instrument Reading x 5)

Sample ID	Type of Analysis	Element	Found / S / I (units)	True / D / SDR (units)	Recalculated		Reported		Acceptable (Y/N)
					%R / RPD / %D	%R / RPD / %D			
D5AB	ICP interference check	Mn	0.4985	0.500	100	NA	NA	Y	
LCS	Laboratory control sample		1.063	1.00	106	106	106	Y	
2	Matrix spike		1.059 (SSR-SR)	1.00	106	106	106	Y	
2/3	Duplicate		1.07	1.059	1	1	1	Y	
1/1	ICP serial dilution								

Comments: Refer to appropriate worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

**Boeing Realty Corp., Bldg C-6 Torrance
Data Validation Reports
LDC# 16739**

Wet Chemistry

LDC

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Boeing Realty Corp., Bldg C-6 Torrance
Collection Date: March 14, 2007
LDC Report Date: May 10, 2007
Matrix: Water
Parameters: Wet Chemistry
Validation Level: Tier 1
Laboratory: TestAmerica

Sample Delivery Group (SDG): IQC1612

Sample Identification

MWB013_WG031407_0001
MWC021_WG031407_0001
MWC021_WG031407_0001MS
MWC021_WG031407_0001MSD

Introduction

This data review covers 4 water samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA Method 300.0 for Chloride, Nitrate, Nitrite, Sulfate, and Orthophosphate, EPA Method 310.1 for Alkalinity, EPA Method 350.3 for Ammonia as Nitrogen, and EPA Method 415.1 for Total Organic Carbon.

The review follows a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Inorganic Data Review (October 2004) as there are no current guidelines for the methods stated above.

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

Blank results are summarized in Section III.

Field duplicates are summarized in Section IX.

Raw data were not reviewed for this SDG. The review was based on QC data.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- N Presumptive evidence of presence of the constituent.
- UU Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. Calibration

a. Initial Calibration

Initial calibration data were not reviewed for Tier 1.

b. Calibration Verification

Calibration verification data were not reviewed for Tier 1.

III. Blanks

Method blanks were reviewed for each matrix as applicable. No contaminant concentrations were found in the preparation blanks.

IV. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) analyses were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

V. Duplicates

Duplicate (DUP) sample analyses were reviewed for each matrix as applicable. Results were within QC limits.

VI. Laboratory Control Samples

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) were within QC limits.

VII. Sample Result Verification

Raw data were not reviewed for this SDG.

VIII. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

IX. Field Duplicates

No field duplicates were identified in this SDG.

X. Field Blanks

No field blanks were identified in this SDG.

**Boeing Realty Corp., Bldg C-6 Torrance
Wet Chemistry - Data Qualification Summary - SDG IQC1612**

No Sample Data Qualified in this SDG

**Boeing Realty Corp., Bldg C-6 Torrance
Wet Chemistry - Laboratory Blank Data Qualification Summary - SDG IQC1612**

No Sample Data Qualified in this SDG

TestAmerica

ANALYTICAL TESTING CORPORATION

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TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727-01
Report Number: IQC1612

Sampled: 03/14/07
Received: 03/14/07

INORGANICS

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC1612-05 (MWB013_WG031407_0001 - Water)									
Reporting Units: mg/l									
Alkalinity as CaCO3	EPA 310.1	7C23075	N/A	2.0	150	1	03/23/07	03/23/07	
Ammonia-N	EPA 350.3	7C26085	N/A	0.50	ND	1	03/26/07	03/26/07	
Chloride	EPA 300.0	7C14053	N/A	25	300	50	03/14/07	03/14/07	
Nitrate-NO3	EPA 300.0	7C14053	N/A	25	60	50	03/14/07	03/14/07	
Nitrite-NO2	EPA 300.0	7C14053	N/A	0.50	ND	1	03/14/07	03/14/07	
Orthophosphate - PO4	EPA 300.0	7C14053	N/A	0.50	ND	1	03/14/07	03/14/07	
Sulfate	EPA 300.0	7C14053	N/A	25	320	50	03/14/07	03/14/07	
Total Organic Carbon	EPA 415.1	7C20118	N/A	1.0	2.1	1	03/20/07	03/20/07	
Sample ID: IQC1612-10 (MWC021_WG031407_0001 - Water)									
Reporting Units: mg/l									
Alkalinity as CaCO3	EPA 310.1	7C23075	N/A	2.0	160	1	03/23/07	03/23/07	
Ammonia-N	EPA 350.3	7C26085	N/A	0.50	ND	1	03/26/07	03/26/07	
Chloride	EPA 300.0	7C14053	N/A	25	120	50	03/14/07	03/14/07	
Nitrate-NO3	EPA 300.0	7C14053	N/A	0.50	9.9	1	03/14/07	03/14/07	
Nitrite-NO2	EPA 300.0	7C14053	N/A	0.50	ND	1	03/14/07	03/14/07	
Orthophosphate - PO4	EPA 300.0	7C14053	N/A	0.50	ND	1	03/14/07	03/14/07	
Sulfate	EPA 300.0	7C14053	N/A	0.50	57	1	03/14/07	03/14/07	
Total Organic Carbon	EPA 415.1	7C20118	N/A	1.0	ND	1	03/20/07	03/20/07	
Sample ID: IQC1612-11 (TMW_10_WG031407_0001 - Water)									
Reporting Units: mg/l									
Alkalinity as CaCO3	EPA 310.1	7C23075	N/A	2.0	280	1	03/23/07	03/23/07	
Ammonia-N	EPA 350.3	7C26085	N/A	0.50	ND	1	03/26/07	03/26/07	
Chloride	EPA 300.0	7C14053	N/A	25	560	50	03/14/07	03/14/07	
Nitrate-NO3	EPA 300.0	7C14053	N/A	25	61	50	03/14/07	03/14/07	
Nitrite-NO2	EPA 300.0	7C14053	N/A	0.50	ND	1	03/14/07	03/14/07	
Orthophosphate - PO4	EPA 300.0	7C14053	N/A	0.50	6.4	1	03/14/07	03/14/07	
Sulfate	EPA 300.0	7C14053	N/A	0.50	48	1	03/14/07	03/14/07	
Total Organic Carbon	EPA 415.1	7C20118	N/A	1.0	1.8	1	03/20/07	03/20/07	

TestAmerica - Irvine, CA
Nicholas Marz
Project Manager

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IQC1612 <Page 27 of 51>

LDC #: 16739A6
 SDG #: IQC1612
 Laboratory: Test America

VALIDATION COMPLETENESS WORKSHEET
 EPA Region 1 - Tier 1

Date: 5/10/07
 Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: Alkalinity (EPA Method 310.1), Ammonia-N (EPA Method 350.3), Chloride, Nitrate-N, Nitrite-N, Sulfate, Orthophosphate-X (EPA Method 300.0), TOC (EPA Method 415.1)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: 3/1/07
IIa.	Initial calibration	N	
IIb.	Calibration verification	N	
III.	Blanks	A	MB
IVa.	Matrix Spike/(Matrix Spike) Duplicates	A	MS/MSD / comp
IVb.	Laboratory control samples	A	LC
V.	Sample result verification	N	
VI.	Overall assessment of data	A	
VII.	Field duplicates	N	
VIII.	Field blanks	N	

Note: A = Acceptable ND = No compounds detected D = Duplicate
 N = Not provided/applicable R = Rinsate TB = Trip blank
 SW = See worksheet FB = Field blank EB = Equipment blank

Validated Samples:

A

1	MWB013_WG031407_0001	11		21		31	
2	MWC021_WG031407_0001	12		22		32	
3	MWC021_WG031407_0001MS	13		23		33	
4	MWC021_WG031407_0001MSD	14		24		34	
5	MB	15		25		35	
6		16		26		36	
7		17		27		37	
8		18		28		38	
9		19		29		39	
10		20		30		40	

Notes: _____

LDC #: 16739 Ab
 SDG #: CUL con

VALIDATION FINDINGS WORKSHEET
Sample Specific Analysis Reference

Page: 1 of 1
 Reviewer: [Signature]
 2nd reviewer: _____

All circled methods are applicable to each sample.

Sample ID	Parameter
1.2	pH TDS (C) F (NO ₃) (NO ₂) (SO ₄) (PO ₄) (ALK) (CN) (NH ₃) TKN (C) TOC (C) CR ⁰⁺ _____
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺ _____
m 2.4	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN (NH ₃) TKN (C) TOC (C) CR ⁰⁺ _____
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺ _____
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺ _____
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺ _____
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺ _____
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺ _____
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺ _____
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺ _____
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺ _____
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺ _____
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺ _____
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺ _____
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺ _____
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺ _____
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺ _____
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺ _____
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺ _____
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺ _____
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺ _____
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺ _____
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺ _____
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺ _____
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺ _____
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺ _____
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺ _____

Comments: _____

LDC Report# 16739B6

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Boeing Realty Corp., Bldg C-6 Torrance
Collection Date: March 22, 2007
LDC Report Date: May 10, 2007
Matrix: Water
Parameters: Wet Chemistry
Validation Level: Tier 1 & 2
Laboratory: TestAmerica

Sample Delivery Group (SDG): IQC2470

Sample Identification

CMW001_WG032207_0001
CMW002_WG032207_0001**
CMW001_WG032207_0001MS
CMW001_WG032207_0001MSD

**Indicates sample underwent Tier 2 review

Introduction

This data review covers one water sample listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA Method 300.0 for Chloride, Nitrate, Nitrite, Sulfate, and Orthophosphate, EPA Method 310.1 for Alkalinity, EPA Method 350.3 for Ammonia as Nitrogen, EPA Method 376.2 for Sulfide, and EPA Method 415.1 for Total Organic Carbon.

The review follows a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Inorganic Data Review (October 2004) as there are no current guidelines for the methods stated above.

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

Blank results are summarized in Section III.

Field duplicates are summarized in Section IX.

Samples indicated by a double asterisk on the front cover underwent a Tier 2 review. A Tier 1 review was performed on all of the other samples. Raw data were not evaluated for the samples reviewed by Tier 1 criteria since this review is based on QC data.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- N Presumptive evidence of presence of the constituent.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. Calibration

a. Initial Calibration

All criteria for the initial calibration of each method were met.

Initial calibration data were not reviewed for Tier 1.

b. Calibration Verification

Calibration verification frequency and analysis criteria were met for each method when applicable.

Calibration verification data were not reviewed for Tier 1.

III. Blanks

Method blanks were reviewed for each matrix as applicable. No contaminant concentrations were found in the initial, continuing and preparation blanks.

IV. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

V. Duplicates

Duplicate (DUP) sample analyses were reviewed for each matrix as applicable. Results were within QC limits.

VI. Laboratory Control Samples

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) were within QC limits.

VII. Sample Result Verification

Raw data were not reviewed for this SDG.

VIII. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

IX. Field Duplicates

No field duplicates were identified in this SDG.

X. Field Blanks

No field blanks were identified in this SDG.

**Boeing Realty Corp., Bldg C-6 Torrance
Wet Chemistry - Data Qualification Summary - SDG IQC2470**

No Sample Data Qualified in this SDG

**Boeing Realty Corp., Bldg C-6 Torrance
Wet Chemistry - Laboratory Blank Data Qualification Summary - SDG IQC2470**

No Sample Data Qualified in this SDG

TestAmerica

ANALYTICAL TESTING CORPORATION

17461 Derian Avenue, Suite 100, Irvine, CA 92614 (949) 261-1022 Fax: (949) 260-3297

TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IQC2470

Sampled: 03/22/07
Received: 03/22/07

INORGANICS

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC2470-08 (MWB003_WG032207_0001 - Water)									
Reporting Units: mg/l									
Alkalinity as CaCO3	EPA 310.1	7D02058	2.0	2.0	340	1	04/02/07	04/02/07	
Ammonia-N	EPA 350.3	7D03070	0.070	0.50	0.089	1	04/03/07	04/03/07	J
Chloride	EPA 300.0	7C23041	5.0	25	460	50	03/23/07	03/24/07	
Nitrate-NO3	EPA 300.0	7C22150	0.25	0.50	8.4	1	03/22/07	03/23/07	
Nitrite-NO2	EPA 300.0	7C22150	3.0	5.0	ND	10	03/22/07	03/23/07	RL1
Orthophosphate - PO4	EPA 300.0	7C22150	0.40	0.50	ND	1	03/22/07	03/23/07	
Sulfate	EPA 300.0	7C22150	0.15	0.50	32	1	03/22/07	03/23/07	
Total Organic Carbon	EPA 415.1	7C29166	0.50	1.0	1.6	1	03/29/07	03/29/07	
Sample ID: IQC2470-10 (WCC06S_WG032207_0001 - Water)									
Reporting Units: mg/l									
Alkalinity as CaCO3	EPA 310.1	7D02058	2.0	2.0	340	1	04/02/07	04/02/07	
Ammonia-N	EPA 350.3	7D03070	0.070	0.50	0.35	1	04/03/07	04/03/07	J
Chloride	EPA 300.0	7C23041	5.0	25	520	50	03/23/07	03/24/07	
Nitrate-NO3	EPA 300.0	7C22150	0.25	0.50	2.9	1	03/22/07	03/23/07	
Nitrite-NO2	EPA 300.0	7C22150	3.0	5.0	ND	10	03/22/07	03/23/07	RL1
Orthophosphate - PO4	EPA 300.0	7C22150	0.40	0.50	ND	1	03/22/07	03/23/07	
Sulfate	EPA 300.0	7C22150	0.15	0.50	18	1	03/22/07	03/23/07	
Total Organic Carbon	EPA 415.1	7C29166	0.50	1.0	5.2	1	03/29/07	03/29/07	
Sample ID: IQC2470-11 (CMW001_WG032207_0001 - Water)									
Reporting Units: mg/l									
Sulfide	EPA 376.2	7C27124	0.010	0.10	0.032	1	03/27/07	03/27/07	J
Sample ID: IQC2470-13 (CMW002_WG032207_0001 - Water)									
Reporting Units: mg/l									
Alkalinity as CaCO3	EPA 310.1	7D02058	2.0	2.0	ND	1	04/02/07	04/02/07	
Ammonia-N	EPA 350.3	7D03070	0.070	0.50	0.17	1	04/03/07	04/03/07	J
Chloride	EPA 300.0	7C22150	1.0	5.0	120	10	03/22/07	03/23/07	
Nitrate-NO3	EPA 300.0	7C22150	0.25	0.50	ND	1	03/22/07	03/23/07	
Nitrite-NO2	EPA 300.0	7C22150	3.0	5.0	ND	10	03/22/07	03/23/07	RL1
Orthophosphate - PO4	EPA 300.0	7C22150	0.40	0.50	ND	1	03/22/07	03/23/07	
Sulfate	EPA 300.0	7C22150	1.5	5.0	98	10	03/22/07	03/23/07	
Sulfide	EPA 376.2	7C27124	0.010	0.10	0.081	1	03/27/07	03/27/07	J
Total Organic Carbon	EPA 415.1	7C29166	0.50	1.0	14	1	03/29/07	03/29/07	

TestAmerica - Irvine, CA
Nicholas Marz
Project Manager

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IQC2470 <Page 52 of 98>

LDC #: 16739B6
 SDG #: IQC2470
 Laboratory: Test America

VALIDATION COMPLETENESS WORKSHEET
 EPA Region 1 - Tier 1/2

Date: 5/10/07
 Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: Alkalinity (EPA Method 310.1), Ammonia-N (EPA Method 350.3), Chloride, Nitrate, Nitrite, Sulfate, Orthophosphate (EPA Method 300.0), Sulfide (EPA Method 376.2), TOC (EPA Method 415.1)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: <u>3/22/07</u>
IIa.	Initial calibration	A	Not reviewed for Tier I validation.
IIb.	Calibration verification	A	Not reviewed for Tier I validation.
III.	Blanks	A	
IVa.	Matrix Spike/(Matrix Spike) Duplicates	A	<u>MS/MSD/MS</u>
IVb.	Laboratory control samples	A	<u>LC</u>
V.	Sample result verification	N	
VI.	Overall assessment of data	A	
VII.	Field duplicates	N	
VIII.	Field blanks	N	

Note: A = Acceptable ND = No compounds detected D = Duplicate
 N = Not provided/applicable R = Rinsate TB = Trip blank
 SW = See worksheet FB = Field blank EB = Equipment blank

Validated Samples: * Indicates sample underwent Tier I validation

1	CMW001_WG032207_0001*	11		21		31	
2	CMW002_WG032207_0001**	12		22		32	
3	CMW001_WG032207_0001MS	13		23		33	
4	CMW001_WG032207_0001MSD	14		24		34	
5	<u>MB</u>	15		25		35	
6		16		26		36	
7		17		27		37	
8		18		28		38	
9		19		29		39	
10		20		30		40	

Notes: _____

LDC #: 16739B6
 SDG #: 502490

VALIDATION FINDINGS WORKSHEET

Sample Specific Analysis Reference

Page: 1 of 1
 Reviewer: W
 2nd reviewer: d

All circled methods are applicable to each sample.

Sample ID	Parameter
1	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺ (S)
2	pH TDS (C) F (NO ₃) (NO ₂) (SO ₄) (PO ₄) ALK CN (NH ₃) TKN (TOC) CR ⁰⁺ (S)
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
3,4	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺ (S)
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺
	pH TDS Cl F NO ₃ NO ₂ SO ₄ PO ₄ ALK CN NH ₃ TKN TOC CR ⁰⁺

Comments: _____

LDC Report# 16739C6

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Boeing Realty Corp., Bldg C-6 Torrance

Collection Date: March 27, 2007

LDC Report Date: May 10, 2007

Matrix: Water

Parameters: Wet Chemistry

Validation Level: Tier 3

Laboratory: TestAmerica, Inc.

Sample Delivery Group (SDG): IQC2895

Sample Identification

MWB019_WG032707_0001

Introduction

This data review covers one water sample listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA Method 300.0 for Chloride, Nitrate, Nitrite, Sulfate, and Orthophosphate, EPA Method 310.1 for Alkalinity, EPA Method 350.3 for Ammonia as Nitrogen, EPA Method 376.2 for Sulfide, and EPA Method 415.1 for Total Organic Carbon.

The review follows a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Inorganic Data Review (October 2004) as there are no current guidelines for the methods stated above.

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

Blank results are summarized in Section III.

Field duplicates are summarized in Section IX.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- N Presumptive evidence of presence of the constituent.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. Calibration

a. Initial Calibration

All criteria for the initial calibration of this method were met.

b. Calibration Verification

Calibration verification frequency and analysis criteria were met for this method when applicable.

III. Blanks

Method blanks were reviewed for each matrix as applicable. No contaminant concentrations were found in the initial, continuing and preparation blanks with the following exceptions:

Method Blank ID	Analyte	Concentration	Associated Samples
MB	Ammonia as N	0.0783 mg/L	All samples in SDG IQC2895
ICB/CCB	Ammonia as N Orthophosphate	0.1038 mg/L 0.436 mg/L	All samples in SDG IQC2895

Sample concentrations were compared to concentrations detected in the method blanks. The sample concentrations were either not detected or were significantly greater (>5X blank contaminants) than the concentrations found in the associated method blanks with the following exceptions:

Sample	Analyte	Reported Concentration	Modified Final Concentration
MWB019_WG032707_0001	Ammonia as N	0.19 mg/L	0.19U mg/L

IV. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) analyses were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

V. Duplicates

Duplicate (DUP) sample analyses were reviewed for each matrix as applicable. Results were within QC limits.

VI. Laboratory Control Samples

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) were within QC limits.

VII. Sample Result Verification

All sample result verifications were acceptable.

VIII. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

IX. Field Duplicates

No field duplicates were identified in this SDG.

X. Field Blanks

No field blanks were identified in this SDG.

**Boeing Realty Corp., Bldg C-6 Torrance
Wet Chemistry - Data Qualification Summary - SDG IQC2895**

No Sample Data Qualified in this SDG

**Boeing Realty Corp., Bldg C-6 Torrance
Wet Chemistry - Laboratory Blank Data Qualification Summary - SDG IQC2895**

SDG	Sample	Analyte	Modified Final Concentration	A or P
IQC2895	MWB019_WG032707_0001	Ammonia as N	0.19U mg/L	A

TestAmerica

ANALYTICAL TESTING CORPORATION

17461 Derian Avenue, Suite 100, Irvine, CA 92614 (949) 261-1022 Fax: (949) 260-3297

TAIT Environmental/Boeing
701 N. Parkcenter Drive
Santa Ana, CA 92705
Attention: Mehmet Pehlivan

Project ID: Boeing C-6 Torrance
EM2727
Report Number: IQC2895

Sampled: 03/27/07
Received: 03/27/07

INORGANICS

Analyte	Method	Batch	MDL Limit	Reporting Limit	Sample Result	Dilution Factor	Date Extracted	Date Analyzed	Data Qualifiers
Sample ID: IQC2895-07 (MWB019_WG032707_0001 - Water)									
Reporting Units: mg/l									
Alkalinity as CaCO3	EPA 310.1	7D05119	2.0	2.0	390	1	04/05/07	04/05/07	
Ammonia-N	EPA 350.3	7D09073	0.070	0.50	0.19 U	1	04/09/07	04/09/07	B, J
Chloride	EPA 300.0	7C27042	2.0	10	290	20	03/27/07	03/28/07	
Nitrate-NO3	EPA 300.0	7C27042	5.0	10	110	20	03/27/07	03/28/07	
Nitrite-NO2	EPA 300.0	7C27042	6.0	10	ND	20	03/27/07	03/28/07	RL1
Orthophosphate - PO4	EPA 300.0	7C27042	0.40	0.50	ND	1	03/27/07	03/27/07	
Sulfate	EPA 300.0	7C27042	3.0	10	560	20	03/27/07	03/28/07	
Sulfide	EPA 376.2	7D02095	0.010	0.10	0.026	1	04/02/07	04/02/07	J
Total Organic Carbon	EPA 415.1	7D03118	0.50	1.0	1.5	1	04/03/07	04/03/07	
Sample ID: IQC2895-08 (MWC017_WG032707_0001 - Water)									
Reporting Units: mg/l									
Alkalinity as CaCO3	EPA 310.1	7D05119	2.0	2.0	200	1	04/05/07	04/05/07	
Ammonia-N	EPA 350.3	7D09073	0.070	0.50	0.12	1	04/09/07	04/09/07	B, J
Chloride	EPA 300.0	7C27042	1.0	5.0	99	10	03/27/07	03/28/07	
Nitrate-NO3	EPA 300.0	7C27042	0.25	0.50	11	1	03/27/07	03/27/07	
Nitrite-NO2	EPA 300.0	7C27042	0.30	0.50	ND	1	03/27/07	03/27/07	
Orthophosphate - PO4	EPA 300.0	7C27042	0.40	0.50	ND	1	03/27/07	03/27/07	
Sulfate	EPA 300.0	7C27042	0.15	0.50	58	1	03/27/07	03/27/07	
Sulfide	EPA 376.2	7D02095	0.010	0.10	0.023	1	04/02/07	04/02/07	J
Total Organic Carbon	EPA 415.1	7D03118	0.50	1.0	ND	1	04/03/07	04/03/07	
Sample ID: IQC2895-09 (IRZMW005_WG032707_0001 - Water)									
Reporting Units: mg/l									
Alkalinity as CaCO3	EPA 310.1	7D05119	2.0	2.0	360	1	04/05/07	04/05/07	
Ammonia-N	EPA 350.3	7D09073	0.070	0.50	3.8	1	04/09/07	04/09/07	
Chloride	EPA 300.0	7C27042	2.0	10	290	20	03/27/07	03/28/07	
Nitrate-NO3	EPA 300.0	7C27042	0.25	0.50	1.5	1	03/27/07	03/27/07	
Nitrite-NO2	EPA 300.0	7C27042	0.30	0.50	ND	1	03/27/07	03/27/07	
Orthophosphate - PO4	EPA 300.0	7C27042	0.40	0.50	ND	1	03/27/07	03/27/07	
Sulfate	EPA 300.0	7C27042	0.15	0.50	30	1	03/27/07	03/27/07	
Sulfide	EPA 376.2	7D02095	0.20	2.0	4.1	20	04/02/07	04/02/07	
Total Organic Carbon	EPA 415.1	7D03118	0.50	1.0	3.2	1	04/03/07	04/03/07	

TestAmerica - Irvine, CA
Nicholas Marz
Project Manager

125107

The results pertain only to the samples tested in the laboratory. This report shall not be reproduced, except in full, without written permission from TestAmerica.

IQC2895 <Page 26 of 76>

LDC #: 16739C6
 SDG #: IQC2895
 Laboratory: Test America

VALIDATION COMPLETENESS WORKSHEET
 EPA Region 1 - Tier 3

Date: 5/10/07
 Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: Alkalinity (EPA Method 310.1), Ammonia-N (EPA Method 350.3), Chloride, Nitrate, Nitrite, Sulfate, Orthophosphate, Sulfide (EPA Method 376.2), TOC (EPA Method 415.1)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: 3/29/07
IIa.	Initial calibration	A	
IIb.	Calibration verification	A	
III.	Blanks	SW	
IVa.	Matrix Spike/(Matrix Spike) Duplicates	A	no direct MS/MSO/MP
IVb.	Laboratory control samples	A	LCs
V.	Sample result verification	A	
VI.	Overall assessment of data	A	
VII.	Field duplicates	N	
VIII.	Field blanks	N	

Note: A = Acceptable ND = No compounds detected D = Duplicate
 N = Not provided/applicable R = Rinsate TB = Trip blank
 SW = See worksheet FB = Field blank EB = Equipment blank

Validated Samples:

1	MWB019_WG032707_0001	11		21		31	
2	MB	12		22		32	
3		13		23		33	
4		14		24		34	
5		15		25		35	
6		16		26		36	
7		17		27		37	
8		18		28		38	
9		19		29		39	
10		20		30		40	

Notes: _____

LDC #: 1693/c6
 SDG #: see cover

VALIDATION FINDINGS CHECKLIST

Page: 1 of 1
 Reviewer: HY
 2nd Reviewer: [Signature]

Method: Inorganics (EPA Method See cover)

Validation Area	Yes	No	NA	Findings/Comments
All technical holding times were met.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Cooler temperature criteria was met.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all instruments calibrated daily, each set-up time?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the proper number of standards used?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all initial calibration correlation coefficients > 0.995?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all initial and continuing calibration verification %Rs within the 90-110% QC limits?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were titrant checks performed as required? (Level IV only)	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were balance checks performed as required? (Level IV only)	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Was a method blank associated with every sample in this SDG?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was there contamination in the method blanks? If yes, please see the Blanks validation completeness worksheet.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were a matrix spike (MS) and duplicate (DUP) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD or MS/DUP. Soil / Water.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the 75-125 QC limits? If the sample concentration exceeded the spike concentration by a factor of 4 or more, no action was taken.	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	(none of next)
Were the MS/MSD or duplicate relative percent differences (RPD) < 20% for waters and < 35% for soil samples? A control limit of < CRDL (< 2X CRDL for soil) was used for samples that were < 5X the CRDL, including when only one of the duplicate sample values were < 5X the CRDL.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was an LCS analyzed for this SDG?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was an LCS analyzed per extraction batch?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the 80-120% (85-115% for Method 300.0) QC limits?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
VI. Regional Quality Assurance and Quality Control				
Were performance evaluation (PE) samples performed?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were the performance evaluation (PE) samples within the acceptance limits?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	

LDC #: 16739 CB
 SDG #: See cover

VALIDATION FINDINGS CHECKLIST

Page: 1 of 1
 Reviewer: WY
 2nd Reviewer: [Signature]

Validation Area	Yes	No	NA	Findings/Comments
Were RLs adjusted to reflect all sample dilutions and dry weight factors applicable to level IV validation?	✓			
Were detection limits < RL?	✓			
Overall assessment of data was found to be acceptable.	✓			
Field duplicate pairs were identified in this SDG.		✓		
Target analytes were detected in the field duplicates.			✓	
Field blanks were identified in this SDG.		✓		
Target analytes were detected in the field blanks.			✓	

LDC #: 1673926
 SDG #: See cover

Page: 1 of 1
 Reviewer: WJ
 2nd Reviewer: RL

VALIDATION FINDINGS WORKSHEET
Level IV Recalculation Worksheet

METHOD: Inorganics, Method See cover

Percent recoveries (%R) for a laboratory control sample and a matrix spike sample were recalculated using the following formula:

$$\%R = \frac{\text{Found}}{\text{True}} \times 100$$
 Where, Found = concentration of each analyte measured in the analysis of the sample. For the matrix spike calculation, Found = SSR (spiked sample result) - SR (sample result).
 True = concentration of each analyte in the source.

A sample and duplicate relative percent difference (RPD) was recalculated using the following formula:

$$RPD = \frac{|S-D|}{(S+D)/2} \times 100$$
 Where, S = Original sample concentration
 D = Duplicate sample concentration

Sample ID	Type of Analyte	Element	Found / S (units)	True / D (units)	Recalculated		Acceptable (Y/N)
					%R / RPD	Reported %R / RPD	
LCS	Laboratory control sample	PO4	4.67	5.00	93	93	Y
10-3-93-36 ↓ IPC-2848-5	Matrix spike sample	NO3-N	2.65 (SSR-SR)	2.00	108	108	Y
	Duplicate sample	NO3-N	2.80	2.80	0	0	Y

Comments: Refer to appropriate worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 16739 cb
 SDG #: See cover

Validatin Findings Worksheet
Initial and Continuing Calibration Calculation Verification

Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

Method: Inorganics, Method See cover

The correlation coefficient (r) for the calibration of Cl was recalculated. Calibration date: 3/20/07

An initial or continuing calibration verification percent recovery (%R) was recalculated for each type of analysis using the following formula:

$\%R = \frac{\text{Found} \times 100}{\text{True}}$
 Where, Found = concentration of each analyte measured in the analysis of the ICV or CCV solution
 True = concentration of each analyte in the ICV or CCV source

Type of analysis	Analyte	Standard	conc. Mg/L	Area	Recalculated		Reported		Acceptable (Y/N)
					r	r ²	r	r ²	
Initial calibration Calibration verification	Cl	s1	0	7197.84	0.999986	0.999986	0.999986	0.999986	Y
		s2	0.2	21609.41					
		s3	0.5	57145.97					
		s4	5	589558.07					
		s5	10	1264022.24					
		s6	20	2754745.98					
		s7	30	4466513.23					
CCV Calibration verification	h ₂	10	0.28		0.83		0.83	0.83	Y
CCV Calibration verification	h ₂	10	0.82		0.88		0.88	0.88	Y
CCV Calibration verification	S	0.342	0.347		0.91		0.91	0.91	Y

Comments: Refer to Calibration Verification findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

**Boeing Realty Corp., Bldg C-6 Torrance
Data Validation Reports
LDC# 16739**

Dissolved Gases

LDC

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Boeing Realty Corp., Bldg C-6 Torrance
Collection Date: March 14, 2007
LDC Report Date: May 10, 2007
Matrix: Water
Parameters: Dissolved Gases
Validation Level: Tier 1
Laboratory: TestAmerica/Air Technology Laboratories, Inc.
Sample Delivery Group (SDG): IQC1612/A7031508

Sample Identification

MWB013_WG031407_0001
MWC021_WG031407_0001

Introduction

This data review covers 2 water samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per Method RSK-175 for Dissolved Gases.

This review follows a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Organic Data Review (October 1999) as there are no current guidelines for the method stated above.

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

Blank results are summarized in Section III.

Field duplicates are summarized in Section IX.

Raw data were not reviewed for this SDG. The review was based on QC data.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- N Presumptive evidence of presence of the constituent.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. Calibration

a. Initial Calibration

Initial calibration data were not reviewed for Tier 1.

b. Calibration Verification

Calibration verification data were not reviewed for Tier 1.

III. Blanks

Method blanks were performed at the required frequency. No dissolved gas contaminants were found in the method blanks.

IV. Accuracy and Precision Data

a. Surrogate Recovery

Not required by the method.

b. Matrix Spike/Matrix Spike Duplicates

The laboratory has indicated that there were no matrix spike (MS) and matrix spike duplicate (MSD) analyses specified for the samples in this SDG, and therefore matrix spike and matrix spike duplicate analyses were not performed for this SDG.

c. Laboratory Control Samples

Laboratory control samples were analyzed at the required frequency. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

V. Target Compound Identification

Raw data were not reviewed for this SDG.

VI. Compound Quantitation and CRQLs

Raw data were not reviewed for this SDG.

VII. System Performance

Raw data were not reviewed for this SDG.

VIII. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

IX. Field Duplicates

No field duplicates were identified in this SDG.

X. Field Blanks

No field blanks were identified in this SDG.

**Boeing Realty Corp., Bldg C-6 Torrance
Dissolved Gases - Data Qualification Summary - SDG IQC1612/A7031508**

No Sample Data Qualified in this SDG

**Boeing Realty Corp., Bldg C-6 Torrance
Dissolved Gases - Laboratory Blank Data Qualification Summary - SDG
IQC1612/A7031508**

No Sample Data Qualified in this SDG

Client: TestAmerica
 Attn: Nicholas Marz

Page 2 of 3
 A7031404

Client's Project: IQC1612
 Date Received: 3/15/2007
 Matrix: Water
 Units: ug/L

Dissolved Gases by EPA Procedure RSKSOP-175										
MW 2013 MWCD21 - W6031407-0001										
Lab No.:	A7031508-01		A7031508-02		A7031508-03		A7031508-04			
Client Sample I.D.:	IQC1612-05		IQC1612-10		IQC1612-11		IQC1612-12			
Date Sampled:	3/14/2007		3/14/2007		3/14/2007		3/14/2007			
Date Analyzed:	3/20/2007		3/20/2007		3/20/2007		3/20/2007			
Analyst Initials:	DT		DT		DT		DT			
Data File:	20mar018		20mar019		20mar020		20mar021			
QC Batch:	070320GC8A1		070320GC8A1		070320GC8A1		070320GC8A1			
Dilution Factor:	1.0		1.0		1.0		1.0			
ANALYTE	PQL	RL	Results	RL	Results	RL	Results	RL	Results	
Methane	1.0	1.0	2.3	1.0	67	1.0	13	1.0	12	
Ethane	2.0	2.0	ND	2.0	ND	2.0	ND	2.0	ND	
Ethylene	3.0	3.0	ND	3.0	ND	3.0	ND	3.0	ND	
Carbon Dioxide	200	200	8,500	200	13,000	200	53,000	200	28,000	
Nitrogen	1,500	1,500	94,000	1,500	94,000	1,500	90,000	1,500	91,000	

PQL = Practical Quantitation Limit
 ND = Not Detected (Below RL)
 RL = PQL X Dilution Factor

Reviewed/Approved By: 
 Mark J. Johnson
 Operations Manager

Date: 3-27-07

The cover letter is an integral part of this analytical report.

1651007



AIRTECHNOLOGY Laboratories, Inc.

18501 E. Gale Avenue, Suite 130 ♦ City of Industry, CA 91748 ♦ Ph: (626) 964-4032 ♦ Fx: (626) 964-5832

LDC #: 16739A51 **VALIDATION COMPLETENESS WORKSHEET**
 SDG #: IQC1612/A7031508 Tier 1
 Laboratory: ~~Del Mar Analytical~~ Air Technology Laboratory, Inc.
Test Analyze

Date: 5/10/07
 Page: 1 of 1
 Reviewer: *[Signature]*
 2nd Reviewer: *[Signature]*

METHOD: GC Dissolved Gases (Method RSK-175)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: 3/14/07
IIa.	Initial calibration	N	
IIb.	Calibration verification	N	
III.	Blanks	A	
IVa.	Surrogate recovery	N	not required
IVb.	Matrix spike/Matrix spike duplicates	N	chem specified
IVc.	Laboratory control samples	A	2cs1D
V.	Target compound identification	N	
VI.	Compound Quantitation and CRQLs	N	
VII.	System Performance	N	
VIII.	Overall assessment of data	A	
IX.	Field duplicates	N	
X.	Field blanks	N	

Note: A = Acceptable ND = No compounds detected D = Duplicate
 N = Not provided/applicable R = Rinsate TB = Trip blank
 SW = See worksheet FB = Field blank EB = Equipment blank

Validated Samples: *water*

S 10	1 ↑	MWB013_WG031407_0001	11	MB - 3/20/07	21		31
	2 ↑	MWC021_WG031407_0001	12		22		32
	3		13		23		33
	4		14		24		34
	5		15		25		35
	6		16		26		36
	7		17		27		37
	8		18		28		38
	9		19		29		39
	10		20		30		40

Notes: _____

LDC Report# 16739B51

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Boeing Realty Corp., Bldg C-6 Torrance
Collection Date: March 22, 2007
LDC Report Date: May 10, 2007
Matrix: Water
Parameters: Dissolved Gases
Validation Level: Tier 2
Laboratory: TestAmerica/Air Technology Laboratory, Inc.

Sample Delivery Group (SDG): IQC2470/A7032601

Sample Identification

CMW002_WG032207_0001

Introduction

This data review covers one water sample listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per Method RSK-175 for Dissolved Gases.

This review follows a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Organic Data Review (October 1999) as there are no current guidelines for the method stated above.

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

Blank results are summarized in Section III.

Field duplicates are summarized in Section IX.

Raw data were not reviewed for this SDG. The review was based on QC data.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- N Presumptive evidence of presence of the constituent.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. Calibration

a. Initial Calibration

Initial calibration of compounds was performed as required by the method.

A curve fit, based on the initial calibration, was established for quantitation. The coefficient of determination (r^2) was greater than or equal to 0.990 .

b. Calibration Verification

Calibration verification was performed at required frequencies. The percent differences (%D) of amounts in continuing standard mixtures were within the 25.0% QC limits.

III. Blanks

Method blanks were performed at the required frequency. No dissolved gas contaminants were found in the method blanks.

IV. Accuracy and Precision Data

a. Surrogate Recovery

Not required by the method.

b. Matrix Spike/Matrix Spike Duplicates

The laboratory has indicated that there were no matrix spike (MS) and matrix spike duplicate (MSD) analyses specified for the samples in this SDG, and therefore matrix spike and matrix spike duplicate analyses were not performed for this SDG.

c. Laboratory Control Samples

Laboratory control samples were analyzed at the required frequency. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

V. Target Compound Identification

Raw data were not reviewed for this SDG.

VI. Compound Quantitation and CRQLs

Raw data were not reviewed for this SDG.

VII. System Performance

Raw data were not reviewed for this SDG.

VIII. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

IX. Field Duplicates

No field duplicates were identified in this SDG.

X. Field Blanks

No field blanks were identified in this SDG.

**Boeing Realty Corp., Bldg C-6 Torrance
Dissolved Gases - Data Qualification Summary - SDG IQC2470/A7032601**

No Sample Data Qualified in this SDG

**Boeing Realty Corp., Bldg C-6 Torrance
Dissolved Gases - Laboratory Blank Data Qualification Summary - SDG
IQC2470/A7032601**

No Sample Data Qualified in this SDG

Client: TestAmerica
 Attn: Nicholas Marz

Page 2 of 3
 A7032601

Client's Project: IQC2470
 Date Received: 3/23/2007
 Matrix: Water
 Units: ug/L

Dissolved Gases by EPA Procedure RSKSOP-175										
CMW002-WG032207-000										
Lab No.:	A7032601-01	A7032601-02	A7032601-03	A7032601-04						
Client Sample ID.:	IQC2470-08	IQC2470-10	IQC2470-13	IQC2470-14						
Date Sampled:	3/22/2007	3/22/2007	3/22/2007	3/22/2007						
Date Analyzed:	3/27/2007	3/27/2007	3/27/2007	3/27/2007						
Analyst Initials:	DT	DT	DT	DT						
Data File:	27mar012	27mar013	27mar014	27mar015						
QC Batch:	070327GCRA1	070327GCRA1	070327GCRA1	070327GCRA1						
Dilution Factor:	1.0	1.0	1.0	1.0						
ANALYTE	PQL	RL	Results	RL	Results	RL	Results	RL	Results	
Methane	1.0	1.0	8.8	1.0	3.4	1.0	1.8	1.0	17,000	
Ethane	2.0	2.0	ND	2.0	ND	2.0	ND	2.0	ND	
Ethylene	3.0	3.0	ND	3.0	ND	3.0	5.9	3.0	12	
Carbon Dioxide	200	200	50,000	200	92,000	200	13,000	200	210,000	
Nitrogen	1,500	1,500	110,000	1,500	110,000	1,500	110,000	1,500	81,000	

PQL = Practical Quantitation Limit
 ND = Not Detected (Below RL)
 RL = PQL X Dilution Factor

Reviewed/Approved By: Mark J. Johnson
 Operations Manager

Date: 4/2/07

The cover letter is an integral part of this analytical report.

HC 57007

LDC #: 16739B51 **VALIDATION COMPLETENESS WORKSHEET**
 SDG #: IQC2470/A7032601 Tier 2
 Laboratory: ~~Del Mar Analytical~~ Air Technology Laboratory, Inc.
 Test America
METHOD: GC Dissolved Gases (Method RSK-175)

Date: 5/10/07
 Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: 3/22/07
IIa.	Initial calibration	Δ	r ² 20.990
IIb.	Calibration verification	Δ	%D ≤ 25
III.	Blanks	Δ	
IVa.	Surrogate recovery	N	not Required
IVb.	Matrix spike/Matrix spike duplicates	N	client specified
IVc.	Laboratory control samples	A	see ID
V.	Target compound identification	N	
VI.	Compound Quantitation and CRQLs	N	
VII.	System Performance	N	
VIII.	Overall assessment of data	Δ	
IX.	Field duplicates	N	
X.	Field blanks	N	

Note: A = Acceptable ND = No compounds detected D = Duplicate
 N = Not provided/applicable R = Rinsate TB = Trip blank
 SW = See worksheet FB = Field blank EB = Equipment blank

Validated Samples: water

1	CMW002_WG032207_0001	11	MB - 3/27/07	21		31	
2		12		22		32	
3		13		23		33	
4		14		24		34	
5		15		25		35	
6		16		26		36	
7		17		27		37	
8		18		28		38	
9		19		29		39	
10		20		30		40	

Notes: _____

LDC Report# 16739C51

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Boeing Realty Corp., Bldg C-6 Torrance
Collection Date: March 27, 2007
LDC Report Date: May 10, 2007
Matrix: Water
Parameters: Dissolved Gases
Validation Level: Tier 3
Laboratory: TestAmerica/Air Technology Laboratory, Inc.
Sample Delivery Group (SDG): IQC2895/A7032807

Sample Identification

MWB019_WG032707_0001

Introduction

This data review covers one water sample listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per Method RSK-175 for Dissolved Gases.

This review follows a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Organic Data Review (October 1999) as there are no current guidelines for the method stated above.

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

Blank results are summarized in Section III.

Field duplicates are summarized in Section IX.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- N Presumptive evidence of presence of the constituent.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. Calibration

a. Initial Calibration

Initial calibration of compounds was performed as required by the method.

A curve fit, based on the initial calibration, was established for quantitation. The coefficient of determination (r^2) was greater than or equal to 0.990 .

b. Calibration Verification

Calibration verification was performed at required frequencies. The percent differences (%D) of amounts in continuing standard mixtures were within the 25.0% QC limits.

III. Blanks

Method blanks were reviewed for each matrix as applicable. No dissolved gas contaminants were found in the method blanks.

IV. Accuracy and Precision Data

a. Surrogate Recovery

Surrogates were not required by the method.

b. Matrix Spike/Matrix Spike Duplicates

The laboratory has indicated that there were no matrix spike (MS) and matrix spike duplicate (MSD) analyses specified for the samples in this SDG, and therefore matrix spike and matrix spike duplicate analyses were not performed for this SDG.

c. Laboratory Control Samples

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

V. Target Compound Identification

All target compound identifications were within validation criteria.

VI. Compound Quantitation and CRQLs

All compound quantitation and CRQLs were within validation criteria.

VII. System Performance

The system performance was acceptable.

VIII. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

IX. Field Duplicates

No field duplicates were identified in this SDG.

X. Field Blanks

No field blanks were identified in this SDG.

**Boeing Realty Corp., Bldg C-6 Torrance
Dissolved Gases - Data Qualification Summary - SDG IQC2895/A7032807**

No Sample Data Qualified in this SDG

**Boeing Realty Corp., Bldg C-6 Torrance
Dissolved Gases - Laboratory Blank Data Qualification Summary - SDG
IQC2895/A7032807**

No Sample Data Qualified in this SDG

Client: TestAmerica
 Attn: Nicholas Marz

Client's Project: IQC2895
 Date Received: 3/28/2007
 Matrix: Water
 Units: ug/L

Dissolved Gases by EPA Procedure RSKSOP-175							
Lab No.:	A7032807-01		A7032807-02		A7032807-03		
Client Sample I.D.:	IQC2895-07		IQC2895-08		IQC2895-09		
Date Sampled:	3/27/2007		3/27/2007		3/27/2007		
Date Analyzed:	4/2/2007		4/2/2007		4/2/2007		
Analyst Initials:	DT		DT		DT		
Data File:	02apr009		02apr010		02apr011		
QC Batch:	070402GC8A1		070402GC8A1		070402GC8A1		
Dilution Factor:	1.0		1.0		1.0		
ANALYTE	PQL	RL	Results	RL	Results	RL	Results
Methane	1.0	1.0	ND	1.0	10,000	1.0	3.5
Ethane	2.0	2.0	ND	2.0	ND	2.0	ND
Ethylene	3.0	3.0	ND	3.0	14	3.0	ND
Carbon Dioxide	200	200	78,000	200	83,000	200	11,000
Nitrogen	1,500	1,500	110,000	1,500	100,000	1,500	110,000

PQL = Practical Quantitation Limit
 ND = Not Detected (Below RL)
 RL = PQL X Dilution Factor

Reviewed/Approved By: Mark J. Johnson
 Mark J. Johnson
 Operations Manager

Date: 4/6/07

The cover letter is an integral part of this analytical report.



AIRTECHNOLOGY Laboratories, Inc.

18501 E. Gale Avenue, Suite 130 ♦ City of Industry, CA 91748 ♦ Ph: (626) 964-4032 ♦ Fx: (626) 964-5832

LDC #: 16739C51 **VALIDATION COMPLETENESS WORKSHEET**
 SDG #: IQC2895/A7032807 Tier 3
 Laboratory: ~~Del Mar Analytical~~/Air Technology Laboratory, Inc.
 Test Amenza
METHOD: GC Dissolved Gases (Method RSK-175)

Date: 5/10/07
 Page: 1 of 1
 Reviewer: B
 2nd Reviewer: A

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	Δ	Sampling dates: 3/27/07
IIa.	Initial calibration	Δ	r ² 20.990
IIb.	Calibration verification	Δ	% D ≤ 25
III.	Blanks	Δ	
IVa.	Surrogate recovery	N	not Required
IVb.	Matrix spike/Matrix spike duplicates	N	client specified
IVc.	Laboratory control samples	A	see ID
V.	Target compound identification	Δ	
VI.	Compound Quantitation and CRQLs	Δ	
VII.	System Performance	Δ	
VIII.	Overall assessment of data	Δ	
IX.	Field duplicates	N	
X.	Field blanks	N	

Note: A = Acceptable ND = No compounds detected D = Duplicate
 N = Not provided/applicable R = Rinsate TB = Trip blank
 SW = See worksheet FB = Field blank EB = Equipment blank

Validated Samples: water

1	MWB019_WG032707_0001	11	MB - 4/2/07	21		31	
2		12		22		32	
3		13		23		33	
4		14		24		34	
5		15		25		35	
6		16		26		36	
7		17		27		37	
8		18		28		38	
9		19		29		39	
10		20		30		40	

Notes: _____

LDC #: 16739CS1
 SDG #: MC cover

VALIDATION FINDINGS CHECKLIST

Page: 1 of 2
 Reviewer: B
 2nd Reviewer: A

Method: GC HPLC

Validation Area	Yes	No	NA	Findings/Comments
III: Technical holding times				
All technical holding times were met.	<input checked="" type="checkbox"/>			
Cooler temperature criteria was met.	<input checked="" type="checkbox"/>			
IV: Initial calibration				
Did the laboratory perform a 5 point calibration prior to sample analysis?	<input checked="" type="checkbox"/>			
Was a linear fit used for evaluation? If yes, were all percent relative standard deviations (%RSD) < 20%?	<input checked="" type="checkbox"/>			
Was a curve fit used for evaluation? If Yes, what was the acceptance criteria used?	<input checked="" type="checkbox"/>			
Did the initial calibration meet the curve fit acceptance criteria?	<input checked="" type="checkbox"/>			
Were the RT windows properly established?	<input checked="" type="checkbox"/>			
IV: Continuing calibration				
What type of continuing calibration calculation was performed? ___%D or %R	<input checked="" type="checkbox"/>			
Was a continuing calibration analyzed daily?	<input checked="" type="checkbox"/>			
Were all percent differences (%D) < 15% or percent recoveries 85-115%?	<input checked="" type="checkbox"/>			
Were all the retention times within the acceptance windows?	<input checked="" type="checkbox"/>			
V: Blanks				
Was a method blank associated with every sample in this SDG?	<input checked="" type="checkbox"/>			
Was a method blank analyzed for each matrix and concentration?	<input checked="" type="checkbox"/>			
Was there contamination in the method blanks? If yes, please see the Blanks validation completeness worksheet.			<input checked="" type="checkbox"/>	
V: Surrogate spikes				
Were all surrogate %R within the QC limits?			<input checked="" type="checkbox"/>	
If the percent recovery (%R) of one or more surrogates was outside QC limits, was a reanalysis performed to confirm %R?			<input checked="" type="checkbox"/>	
If any %R was less than 10 percent, was a reanalysis performed to confirm %R?			<input checked="" type="checkbox"/>	
VII: Matrix spike/Matrix spike duplicates				
Were a matrix spike (MS) and matrix spike duplicate (MSD) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD. Soil / Water.			<input checked="" type="checkbox"/>	
Was a MS/MSD analyzed every 20 samples of each matrix?			<input checked="" type="checkbox"/>	
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the QC limits?			<input checked="" type="checkbox"/>	
VIII: Laboratory controls samples				
Was an LCS analyzed for this SDG?	<input checked="" type="checkbox"/>			
Was an LCS analyzed per extraction batch?	<input checked="" type="checkbox"/>			

LDC #: 16739C51
 SDG #: pt cover

VALIDATION FINDINGS CHECKLIST

Page: 2 of 2
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

Validation Area	Yes	No	NA	Findings/Comments
IX Regional Quality Assurance and Quality Control Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the QC limits?	/			
IXa Performance evaluation (PE) samples performed?			/	
IXb Target compound identification Were the performance evaluation (PE) samples within the acceptance limits?				
X Compound quantitation/CRQLs Were the retention times of reported detects within the RT windows?	/			
XI System performance Were compound quantitation and CRQLs adjusted to reflect all sample dilutions and dry weight factors applicable to level IV validation?	/			
XII Overall assessment of data System performance was found to be acceptable.	/			
XIII Field duplicates Overall assessment of data was found to be acceptable.	/			
XIV Field blanks Were field duplicate pairs identified in this SDG?		/		
XV Field blanks Were target compounds detected in the field duplicates?			/	
Were field blanks identified in this SDG?		/		
Were target compounds detected in the field blanks?			/	

LDC # 16739CS/
 SDG# per cover

VALIDATION FINDINGS WORKSHEET
Initial Calibration Calculation Verification

Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: GC Carbon Dioxide (Method RSK-175)

Parameter: Methane Carbon Dioxide

Order of regression: 1

Date	Column/Detector	Compound	X Mass (ppmV)	Y Area
05/24/2007	TCD	CO2	100	552
			1000	3718
	Front		5000	18595
			10000	40200
			100000	428338
			500000	1987935.0

Regression Output:		Reported
Constant	0.0	0.0
Std Err of Y Est	13500.069	
R Squared	0.99971	0.999749
No. of Observations	6.000	
Degrees of Freedom	5.000	
X Coefficient(s)	3.98768E+000	3.98770E+000
Std Err of Coef.	0.03	

LDC #: 16739c51
 SDG #: per count

VALIDATION FINDINGS WORKSHEET
Continuing Calibration Results Verification

Page: 1 of 1
 Reviewer: PK
 2nd Reviewer: PK

METHOD: GC HPLC

The percent difference (%D) of the initial calibration average Calibration Factors (CF) and the continuing calibration CF were recalculated for the compounds identified below using the following calculation:

% Difference = $100 \cdot (\text{ave. CF} - \text{CF}) / \text{ave. CF}$ Where: ave. CF = initial calibration average CF
 CF = A/C
 A = Area of compound
 C = Concentration of compound

#	Standard ID	Calibration Date	Compound	Average CF(ical)/ CCV Conc.	Reported		Recalculated	
					CF/Conc. CCV	%R	CF/Conc. CCV	%D
1	cew 9:09	4/2/07	CO ₂	10000	12151	21.5	12151	21.5
2								
3								
4								

Comments: Refer to Continuing Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 16739CS /
 SDG #: per count

VALIDATION FINDINGS WORKSHEET
Laboratory Control Sample/Laboratory Control Sample Duplicates Results Verification

Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: GC HPLC

The percent recoveries (%R) and relative percent differences (RPD) of the laboratory control sample and laboratory control sample duplicate were recalculated for the compounds identified below using the following calculation:

%Recovery = $100 * ((SSC - SC) / SA)$ Where SSC = Spiked sample concentration SC = Sample concentration SA = Spike added
 RPD = $((SSCLCS - SSCLCSD) * 2) / ((SSCLCS + SSCLCSD)) * 100$ LCS = Laboratory Control Sample LCSD = Laboratory Control Sample duplicate

LCS/LCSD samples: Les 1P

Compound	Spike Added (ppm)		Sample Conc. (ppm)	Spike Sample Concentration (ppm)		LCS Percent Recovery		LCSD Percent Recovery		LCS/LCSD RPD	
	LCS	LCSD		LCS	LCSD	Reported	Recalc.	Reported	Recalc.	Reported	Recalc.
Gasoline (8015)											
Diesel (8015)											
Benzene (8021B)											
Methane (RSK-175)	7000	7000	0	7404.3	6375.8	108	108	92	91.1	16	16
2,4-D (8151)											
Dinoseb (8151)											
Naphthalene (8310)											
Anthracene (8310)											
HMX (8330)											
2,4,6-Trinitrotoluene (8330)											

Comments: Refer to Laboratory Control Sample/Laboratory Control Sample Duplicate findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

VALIDATION FINDINGS WORKSHEET
Sample Calculation Verification

LDC #: 16739 CS1
 SDG #: per contract

METHOD: GC HPLC

Were all reported results recalculated and verified for all level IV samples?
 Were all recalculated results for detected target compounds agree within 10% of the reported results?

Y N / N/A
Y N / N/A

Concentration = $\frac{(A)(Fv)(Df)}{(RF)(Vs \text{ or } Ws)(\%S/100)}$
 Example: Sample ID: #1 Compound Name: CO2

Concentration = _____
 A= Area or height of the compound to be measured
 Fv= Final Volume of extract
 Df= Dilution Factor
 RF= Average response factor of the compound in the initial calibration
 Vs= Initial volume of the sample
 Ws= Initial weight of the sample
 %S= Percent Solid

$y = mx$
 $183283 = (3.9877)(x)$
 $x = 45962.258 = 0.04596 \text{ ppm}$

#	Sample ID	Compound	Reported Concentrations	Recalculated Results Concentrations	Qualifications
		gas in HS =	$0.04596(55.5)(44)(100) = 1640$	68.44	
		gas in liquid =	$0.04596(4)(100)(273) = 9.18 \text{ mg/L}$ $(22.4)(36)(298)$		
		TOTAL =	$(68.44 + 9.18)(1600) = 78,000 \text{ ug/l}$		

Comments: _____

